



STIC Search Report

Biotech-Chem Library

STIC Database Tracking Number: 113864

TO: Hong Liu
Location: REM-5C11
Art Unit: 1624
Wednesday, February 11, 2004
Case Serial Number: 10/075847

From: Peggy Ruppel
Location: Biotech-Chem Library
Phone: 571-272-2557
REM E01b65
peggy.ruppel@uspto.gov

Search Notes

Dear Examiner Liu:

Please see attached results. Barb O'Bryen supervised my work on this search as part of my training, so she did, indeed, work on this search.

Feel free to contact me if you have any questions.

Thank you for using STIC services

Peggy Ruppel
~~308-2270~~ 272-2557

=> b reg

FILE 'REGISTRY' ENTERED AT 11:27:54 ON 11 FEB 2004
 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.
 PLEASE SEE "HELP USAGETERMS" FOR DETAILS.
 COPYRIGHT (C) 2004 American Chemical Society (ACS)

Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

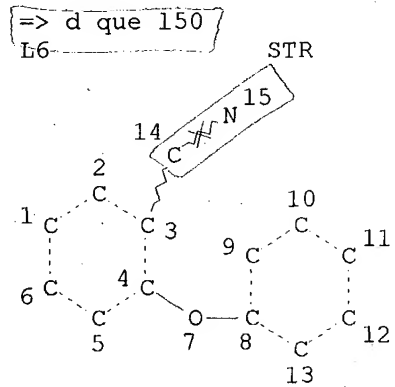
STRUCTURE FILE UPDATES: 10 FEB 2004 HIGHEST RN 648858-13-3
 DICTIONARY FILE UPDATES: 10 FEB 2004 HIGHEST RN 648858-13-3

TSCA INFORMATION NOW CURRENT THROUGH JULY 14, 2003

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. For more information enter HELP PROP at an arrow prompt in the file or refer to the file summary sheet on the web at:
<http://www.cas.org/ONLINE/DBSS/registryss.html>



This structure represents our parent compound for the search. The highlighted nodes and bond are designated as ring or chain throughout the following structures.

NODE ATTRIBUTES:

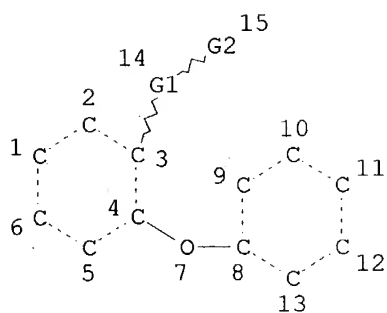
NSPEC IS RC AT 14
 NSPEC IS RC AT 15
 DEFAULT MLEVEL IS ATOM
 DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED
 NUMBER OF NODES IS 15

STEREO ATTRIBUTES: NONE

L7 STR

NH~Ak
@16 17Ak~^N~Ak
18 @19 20CH~Ak
@21 22Ak~^C~^Ak
23 @24 25CH~Ak~^X
@26 27 28Ak~^C~^Ak~^X
29 @30 31 32X~^Ak~^C~^Ak~^X
33 34 @35 36 37

VAR G1=21/24/26/30/35/CH2

VAR G2=16/19/NH2

NODE ATTRIBUTES:

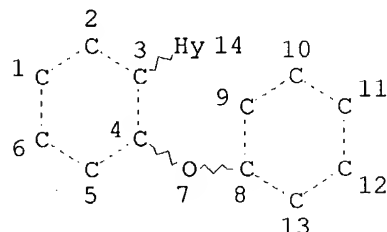
CONNECT IS E1 RC AT 17
 CONNECT IS E1 RC AT 18
 CONNECT IS E1 RC AT 20
 CONNECT IS E1 RC AT 22
 CONNECT IS E1 RC AT 23
 CONNECT IS E1 RC AT 25
 CONNECT IS E1 RC AT 29
 DEFAULT MLEVEL IS ATOM
 DEFAULT ECLEVEL IS LIMITED
 ECOUNT IS M1-X4 C AT 17
 ECOUNT IS M1-X4 C AT 18
 ECOUNT IS M1-X4 C AT 20
 ECOUNT IS M1-X4 C AT 22
 ECOUNT IS M1-X4 C AT 23
 ECOUNT IS M1-X4 C AT 25

GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED
 NUMBER OF NODES IS 37

STEREO ATTRIBUTES: NONE

L8 STR



NODE ATTRIBUTES:

DEFAULT MLEVEL IS ATOM
 GGCAT IS MCY SAT AT 14

Structures L7 and L8
 are two general sub-
 structures of L6 and
 were used to narrow
 the search.

Heterocycle at node 14 is monocyclic and
 unsaturated.

DEFAULT ECLEVEL IS LIMITED
ECOUNT IS M1 N AT 14

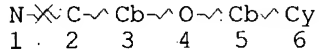
GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED
NUMBER OF NODES IS 14

STEREO ATTRIBUTES: NONE

L9 (6520) SEA FILE=REGISTRY SSS FUL L6

L10 (747) SEA FILE=REGISTRY SUB=L9 SSS FUL (L6 AND (L7 OR L8))
L11 STR



structures L11-L14
were known to be
unacceptable and were
removed from the
answerset.

(Cyclic groups off of
phenyl/naphthyl groups)

NODE ATTRIBUTES:

NSPEC IS RC AT 1

NSPEC IS RC AT 2

DEFAULT MLEVEL IS ATOM

DEFAULT ECLEVEL IS LIMITED

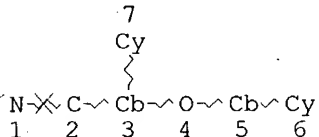
GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED

NUMBER OF NODES IS 6

STEREO ATTRIBUTES: NONE

L12 STR



NODE ATTRIBUTES:

NSPEC IS RC AT 1

NSPEC IS RC AT 2

DEFAULT MLEVEL IS ATOM

DEFAULT ECLEVEL IS LIMITED

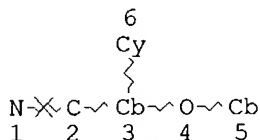
GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED

NUMBER OF NODES IS 7

STEREO ATTRIBUTES: NONE

L13 STR



NODE ATTRIBUTES:

NSPEC IS RC AT 1
 NSPEC IS RC AT 2
 DEFAULT MLEVEL IS ATOM
 DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED
 NUMBER OF NODES IS 6

STEREO ATTRIBUTES: NONE
 L14 STR

N->C~Cb~O~Cb
 1 2 3 4 5

NODE ATTRIBUTES:

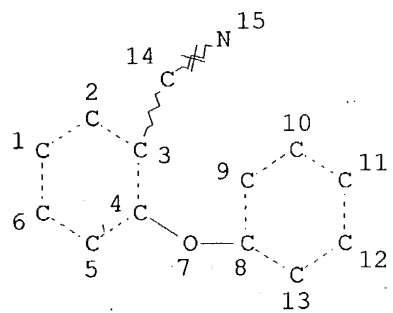
NSPEC IS RC AT 1
 NSPEC IS RC AT 2
 DEFAULT MLEVEL IS ATOM
 DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED
 NUMBER OF NODES IS 5

STEREO ATTRIBUTES: NONE

L15 544 SEA FILE=REGISTRY SUB=L10 SSS FUL (L6 AND L14 NOT (L11 OR L12
 OR L13)).
 L17 STR



this is a restatement of L6

NODE ATTRIBUTES:
 NSPEC IS RC AT 14
 NSPEC IS RC AT 15
 DEFAULT MLEVEL IS ATOM
 DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:
 RING(S) ARE ISOLATED OR EMBEDDED
 NUMBER OF NODES IS 15

STEREO ATTRIBUTES: NONE
 L20 STR

structures L20, and
 L46-L48 were used to
 finalize the search

N~X C~ Cb~ O~ Cb
 1 2 3 4 5

NODE ATTRIBUTES:

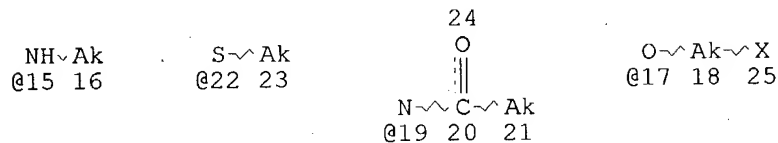
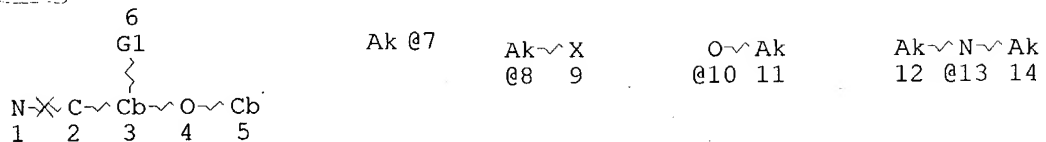
NSPEC IS RC AT 1
 NSPEC IS RC AT 2
 CONNECT IS E2 RC AT 3
 CONNECT IS E1 RC AT 5
 DEFAULT MLEVEL IS ATOM
 DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED
 NUMBER OF NODES IS 5

STEREO ATTRIBUTES: NONE

L46 } STR



VAR G1=X/7/8/10/13/15/22/19/22/17/CN/NO2/NH2

NODE ATTRIBUTES:

NSPEC IS RC AT 1
 NSPEC IS RC AT 2
 CONNECT IS E1 RC AT 5
 CONNECT IS E1 RC AT 7
 CONNECT IS E1 RC AT 11
 CONNECT IS E1 RC AT 12
 CONNECT IS E1 RC AT 14
 CONNECT IS E1 RC AT 16
 CONNECT IS E1 RC AT 21
 CONNECT IS E1 RC AT 23
 DEFAULT MLEVEL IS ATOM
 DEFAULT ECLEVEL IS LIMITED

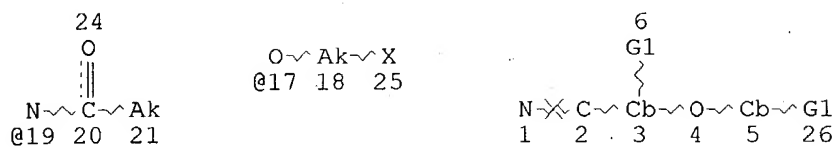
GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED
 NUMBER OF NODES IS 25

STEREO ATTRIBUTES: NONE

L47 } STR

Ak @7	Ak~X @8 9	O~Ak @10 11	Ak~N~Ak 12 @13 14	NH~Ak @15 16	S~Ak @22 23
-------	--------------	----------------	----------------------	-----------------	----------------



VAR G1=X/7/8/10/13/15/22/19/22/17/CN/NO2/NH2

NODE ATTRIBUTES:

```

NSPEC IS RC AT 1
NSPEC IS RC AT 2
CONNECT IS E1 RC AT 7
CONNECT IS E1 RC AT 11
CONNECT IS E1 RC AT 12
CONNECT IS E1 RC AT 14
CONNECT IS E1 RC AT 16
CONNECT IS E1 RC AT 21
CONNECT IS E1 RC AT 23
DEFAULT MLEVEL IS ATOM
DEFAULT ECLEVEL IS LIMITED
  
```

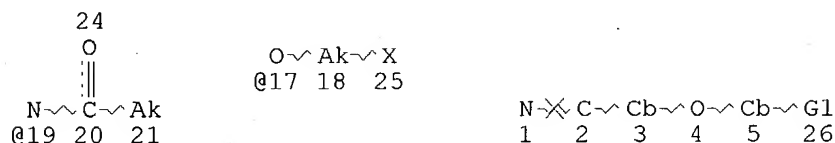
GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED
 NUMBER OF NODES IS 26

STEREO ATTRIBUTES: NONE

{ L48 } STR

Ak @7	Ak~X @8 9	O~Ak @10 11	Ak~N~Ak 12 @13 14	NH~Ak @15 16	S~Ak @22 23
-------	--------------	----------------	----------------------	-----------------	----------------



VAR G1=X/7/8/10/13/15/22/19/22/17/CN/NO2/NH2

NODE ATTRIBUTES:

```

NSPEC IS RC AT 1
NSPEC IS RC AT 2
CONNECT IS E2 RC AT 3
CONNECT IS E1 RC AT 7
CONNECT IS E1 RC AT 11
CONNECT IS E1 RC AT 12
CONNECT IS E1 RC AT 14
CONNECT IS E1 RC AT 16
CONNECT IS E1 RC AT 21
CONNECT IS E1 RC AT 23
DEFAULT MLEVEL IS ATOM
  
```

DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED
NUMBER OF NODES IS 25

STEREO ATTRIBUTES: NONE

L50 } 273 SEA FILE=REGISTRY SUB=L15 SSS FUL (L17 AND (L20 OR L46 OR L47
OR L48)) }

=> b hcaplus

{FILE 'HCAPLUS' ENTERED AT 11:28:09 ON 11 FEB 2004
USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.
PLEASE SEE "HELP USAGETERMS" FOR DETAILS.
COPYRIGHT (C) 2004 AMERICAN CHEMICAL SOCIETY (ACS)

Copyright of the articles to which records in this database refer is held by the publishers listed in the PUBLISHER (PB) field (available for records published or updated in Chemical Abstracts after December 26, 1996), unless otherwise indicated in the original publications. The CA Lexicon is the copyrighted intellectual property of the American Chemical Society and is provided to assist you in searching databases on STN. Any dissemination, distribution, copying, or storing of this information, without the prior written consent of CAS, is strictly prohibited.

FILE COVERS 1907 - 11 Feb 2004 VOL 140 ISS 7
FILE LAST UPDATED: 10 Feb 2004 (20040210/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

'OBI' IS DEFAULT SEARCH FIELD FOR 'HCAPLUS' FILE

=> d que 151 nos }

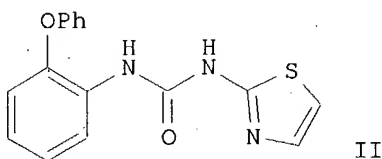
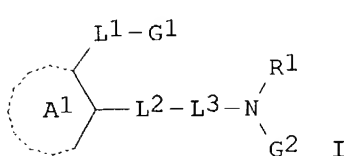
L6	STR
L7	STR
L8	STR
L9	(6520) SEA FILE=REGISTRY SSS FUL L6
L10	(747) SEA FILE=REGISTRY SUB=L9 SSS FUL (L6 AND (L7 OR L8))
L11	STR
L12	STR
L13	STR
L14	STR
L15	544 SEA FILE=REGISTRY SUB=L10 SSS FUL (L6 AND L14 NOT (L11 OR L12 OR L13))
L17	STR
L20	STR
L46	STR
L47	STR
L48	STR
L50	273 SEA FILE=REGISTRY SUB=L15 SSS FUL (L17 AND (L20 OR L46 OR L47 OR L48))
L51	44 SEA FILE=HCAPLUS ABB=ON PLU=ON L50 }

{=> d ibib abs hitstr 151 1-44 }

51 ANSWER 1 OF 44 HCAPLUS COPYRIGHT 2004 ACS on STN
 ACCESSION NUMBER: 2004:20494 HCAPLUS
 DOCUMENT NUMBER: 140:77140
 TITLE: Preparation of thiazolyl aryl ureas as activators of glucokinase
 INVENTOR(S): Polisetti, Dharma Rao; Kodra, Janos Tibor; Lau,
 Jesper; Bloch, Paw; Valcarce-Lopez, Maria Carmen;
 Blume, Niels; Guzel, Mustafa; Santhosh, Kalpathy
 Chidambareswaran; Mjalli, Adnan M. M.; Andrews, Robert
 Carl; Subramanian, Govindan; Ankersen, Michael; Vedso,
 Per; Murray, Anthony; Jeppesen, Lone
 PATENT ASSIGNEE(S): Novo Nordisk A/S, Den.; Valcarce-Lopez, mariacarmen; et al.
 SOURCE: PCT Int. Appl., 600 pp.
 CODEN: PIXXD2
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 9
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2004002481	A1	20040108	WO 2003-DK449	20030627
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU				
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
PRIORITY APPLN. INFO.:			DK 2002-999	A 20020627
			US 2002-394144P	P 20020703
			DK 2003-286	A 20030225
			US 2003-452228P	P 20030305

GI



AB The title compds. [I; A1 = arylene, heteroarylene, fused cycloalkylarylene, etc.; L1 = a bond, O, S, SO, etc.; G1 = alkyl, cycloalkyl, cycloalkylalkylene, etc.; L2 = a bond, alkylene, alkenylene,

etc.; L3 = CO, COCO, COCH₂CO, SO₂; R1 = alkyl, alkenyl, alkynyl, etc.; G2 = heteroaryl, fused heterocyclylheteroaryl, cycloalkylheteroaryl, etc.] which are activators of glucokinase and may be useful for the management, treatment, control, or adjunct treatment of diseases, where increasing glucokinase activity is beneficial (no data), were prepared and formulated. Thus, reacting 2-phenoxyaniline with 2-aminothiazole and 1,1'-carbonyldimidazole afforded 95% the urea II.

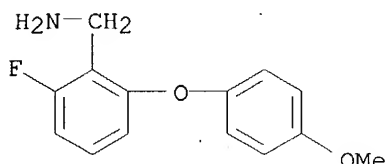
IT 361394-40-3, 2-Fluoro-6-(4-methoxyphenoxy)benzylamine

RL: RCT (Reactant); RACT (Reactant or reagent)

(preparation of thiazolyl aryl ureas as glucokinase activators)

RN 361394-40-3 HCAPLUS

CN Benzenemethanamine, 2-fluoro-6-(4-methoxyphenoxy)- (9CI) (CA INDEX NAME)



REFERENCE COUNT: 20 THERE ARE 20 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

~~D51~~ ANSWER 2 OF 44 HCAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: 2003:777769 HCAPLUS

DOCUMENT NUMBER: 139:261302

TITLE: Preparation of aminobenzamide derivatives for treatment of diabetes and obesity

INVENTOR(S): Nishimura, Teruyuki; Iino, Tomoharu; Nagata, Yasufumi; Eiki, Junichi

PATENT ASSIGNEE(S): Banyu Pharmaceutical Co., Ltd., Japan

SOURCE: PCT Int. Appl., 112 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

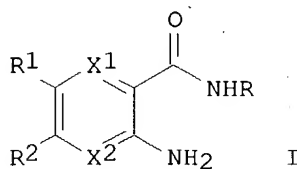
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2003080585	A1	20031002	WO 2003-JP3656	20030325
W:	AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM			
RW:	GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG			
PRIORITY APPLN. INFO.:			JP 2002-85720	A 20020326
OTHER SOURCE(S):		MARPAT 139:261302		

GI



AB Title compds. I (R = nitrogen-containing, monocyclic or bicyclic heteroaryl group; R1 = heterocyclylthio, heterocyclylsulfinyl, heterocyclylsulfonyl, etc.; R2 = H, halo, alkyl, etc.; X1, X2 = N, CH) and their pharmaceutically acceptable salts, having glucokinase activity and useful for prevention or treatment of diabetes and obesity, are prepared. 2-Amino-5-(4-methyl-4H-1,2,4-triazol-3-ylthio)-N-(4-methylthiazol-2-yl)benzamide was prepared and showed hypoglycemic activity in mice.

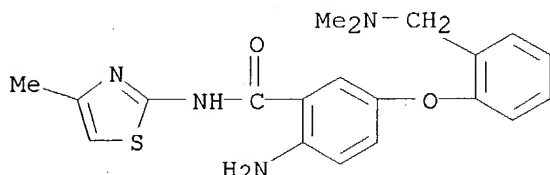
IT 603109-35-9P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of aminobenzamide derivs. for treatment of diabetes and obesity)

RN 603109-35-9 HCPLUS

CN Benzamide, 2-amino-5-[2-[(dimethylamino)methyl]phenoxy]-N-(4-methyl-2-thiazolyl)-(9CI) (CA INDEX NAME)



REFERENCE COUNT: 17 THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L51 ANSWER 3 OF 44 HCPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: 2003:696859 HCPLUS

DOCUMENT NUMBER: 139:230480

TITLE: Preparation of substituted amines prodrugs useful in treating Alzheimer's disease

INVENTOR(S): Varghese, John; Jagodzinska, Barbara; Maillard, Michel; Beck, James P.; Tenbrink, Ruth E.; Getman, Daniel

PATENT ASSIGNEE(S): Elan Pharmaceuticals, Inc., USA; Pharmacia & Upjohn

SOURCE: PCT Int. Appl., 483 pp.

CODEN: PIXXD2

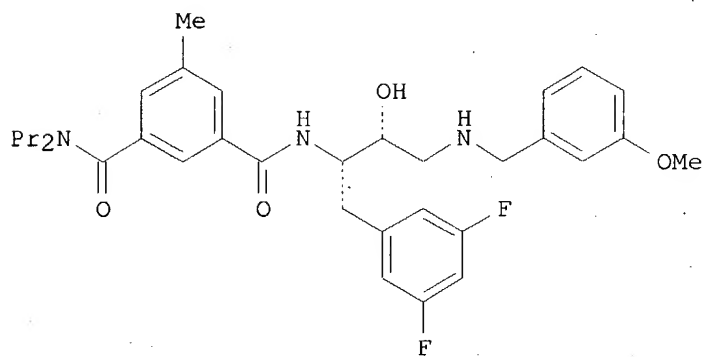
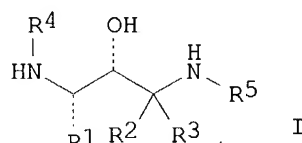
DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2003072535	A2	20030904	WO 2003-US7287	20030227
W:	AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM			
RW:	GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG			
PRIORITY APPLN. INFO.:			US 2002-359953P	P 20020227
OTHER SOURCE(S):		MARPAT 139:230480		



AB Amines [I; R₁ = (un)substituted alkyl, alkenyl, alkynyl, etc.; R₂ = H, (un)substituted alkyl, alkenyl, etc.; R₃ = H, (un)substituted alkyl, alkenyl, etc.; R₄ = XR; X = CO, SO₂, a bond, etc.; R = Ph, naphthyl, indanyl, etc.; R₅ = (un)substituted alkyl, (CH₂)₀₋₃cycloalkyl, etc.; e.g. N1-[(1S,2R)-1-(3,5-difluorobenzyl)-2-hydroxy-3-[(3-methoxybenzyl)amino]propyl]-5-methyl-N3,N3-dipropylisophthalamide], useful in treating Alzheimer's disease and other similar diseases, were prepared. Although the methods of preparation are not claimed, hundreds of example preps. are included. Thus, reacting (2R,3S)-3-amino-4-(3,5-difluorophenyl)-1-[(3-methoxybenzyl)amino]-2-butanol trifluoroacetate with 5-methyl-N,N-dipropylisophthalamic acid in the presence of Et₃N,

1-hydroxybenzotriazole and 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride in DMF afforded (1S,2R)-II (N1-[(1S,2R)-1-(3,5-difluorobenzyl)-2-hydroxy-3-[(3-methoxybenzyl)amino]propyl]-5-methyl-N3,N3-dipropylisophthalamide). The compds. I exhibit an IC₅₀ of < 50 μM against β-secretase.

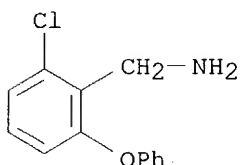
IT 175136-89-7, [(2-Chloro-6-phenoxyphenyl)methyl]amine

RL: RCT (Reactant); RACT (Reactant or reagent)

(preparation of substituted amine prodrugs useful in treating Alzheimer's disease)

RN 175136-89-7 HCPLUS

CN Benzenemethanamine, 2-chloro-6-phenoxy- (9CI) (CA INDEX NAME)



ANSWER 4 OF 44 HCPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: 2003:678291 HCPLUS

DOCUMENT NUMBER: 139:202503

TITLE: Osmotic delivery system containing a polyethylene oxide and an osmagent

INVENTOR(S): Waterman, Kenneth C.

PATENT ASSIGNEE(S): USA

SOURCE: U.S. Pat. Appl. Publ., 12 pp.

CODEN: USXXCO

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2003161882	A1	20030828	US 2003-352258	20030127

PRIORITY APPLN. INFO.: US 2002-353502P P 20020201

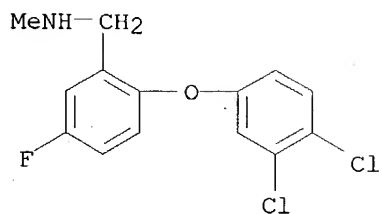
AB An osmotic pharmaceutical tablet is described which comprises a single-layer compressed core surrounded by a water permeable layer having a passageway. The single-layer core contains (i) a non-ripening drug having a solubility per dose less than about 1 mL -1 , (ii) about 2.0% to about 30% by weight of a polyethylene oxide having a weight-average, mol. weight from about 200,000 to about 7,000,000, (iii) an osmagent, and (iv) an optional disintegrant. Many osmotic tablets were prepared and their dissoln. rate were studied.

IT 289716-93-4

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(osmotic delivery system containing polyethylene oxide and osmagent)

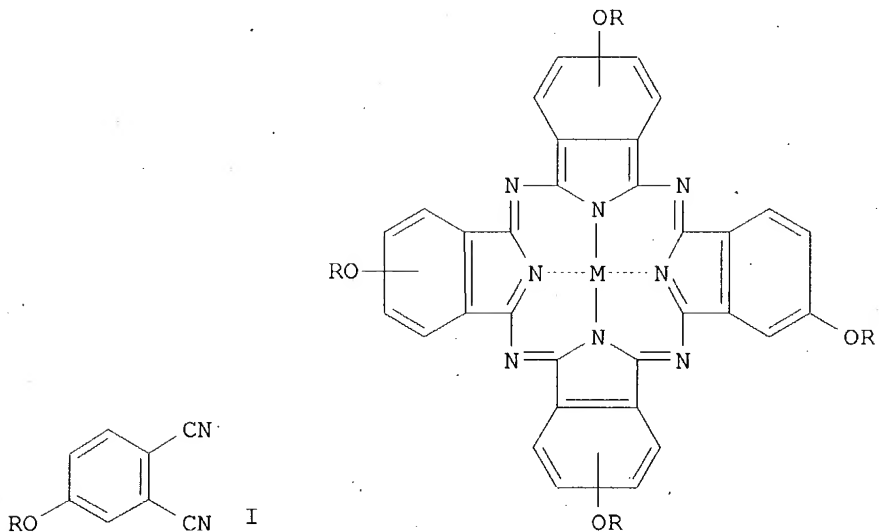
RN 289716-93-4 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N-methyl-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

~~DS1~~ ANSWER 5 OF 44 HCPLUS COPYRIGHT 2004 ACS on STN
 ACCESSION NUMBER: 2003:416216 HCPLUS
 DOCUMENT NUMBER: 139:337765
 TITLE: Microwave-assisted synthesis of phthalonitriles and phthalocyanines
 AUTHOR(S): Csokai, Viktor; Parlagh, Gyula; Grofcsik, Andras;
 Kubinyi, Miklos; Bitter, Istvan
 CORPORATE SOURCE: Department of Organic Chemical Technology, Budapest
 University of Technology and Economics, Budapest,
 H-1521, Hung.
 SOURCE: Synthetic Communications (2003), 33(10), 1615-1621
 CODEN: SYNCV; ISSN: 0039-7911
 PUBLISHER: Marcel Dekker, Inc.
 DOCUMENT TYPE: Journal
 LANGUAGE: English
 GI



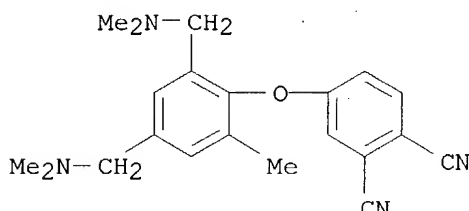
AB The microwave-assisted synthesis of phthalonitriles such as I [R = Me(OCH₂CH₂)₂] and phthalocyanines such as II (same R; M = Zn, 2H) using a two-step literature method was compared with that of a one pot procedure. The two step literature method, comprising first the preparation of the appropriate phthalonitrile followed by cyclization, generally used for the synthesis of PCs was affected by microwave (Mw) irradiation. The synthetic route was based on the well known nucleophilic substitution of 4-nitrophthalonitrile with alcs. and phenols in the presence of K₂CO₃ in DMF solvent. The one pot procedure provides remarkable advantages compared with the two step synthesis of PC-Zn complexes: the overall yields are comparable, the expensive DBU can be eliminated, the work-up is easier, and this method is suitable for direct preparation of metal-free PCs.

IT 615249-96-2P

RL: SPN (Synthetic preparation); PREP (Preparation)
(microwave-assisted synthesis of phthalonitriles and phthalocyanines)

RN 615249-96-2 HCAPLUS

CN 1,2-Benzenedicarbonitrile, 4-[2,4-bis[(dimethylamino)methyl]-6-methylphenoxy]- (9CI) (CA INDEX NAME)



REFERENCE COUNT: 22 THERE ARE 22 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

~~✓~~ ANSWER 6 OF 44 HCAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: 2003:289083 HCAPLUS

DOCUMENT NUMBER: 139:360973

TITLE: Molecular modeling of potential new and selective PET radiotracers for the serotonin transporter

AUTHOR(S): Wellsow, Julia; Kovar, Karl-Artur; Machulla, Hans-Jurgen

CORPORATE SOURCE: Pharmaceutical Institute, Department of Pharmaceutical and Analytical Chemistry, University of Tubingen, Tubingen, Germany

SOURCE: Journal of Pharmacy & Pharmaceutical Sciences [online computer file] (2002), 5(3), 245-257

CODEN: JPPSFY; ISSN: 1482-1826

URL: [http://www.ualberta.ca/~csp/JPPS5\(3\)/K.Kovar/pet.pdf](http://www.ualberta.ca/~csp/JPPS5(3)/K.Kovar/pet.pdf)

PUBLISHER: Canadian Society for Pharmaceutical Sciences

DOCUMENT TYPE: Journal; (online computer file)

LANGUAGE: English

AB Imaging the serotonin transporter (SERT) with Positron Emission Tomography (PET) provides a useful tool for understanding alterations of the serotonergic system. However, no optimal PET radiotracer for the SERT yet exists. The main purpose of this study was to design potential new and

selective PET radiotracers for the SERT and to predict their binding affinity at both the SERT and the norepinephrine transporter. Mol. Modeling was used for ligand design. Predictions of binding affinity were based on models generated by Comparative Mol. Field Anal. (CoMFA) and Comparative Mol. Similarity Indexes Anal. (CoMSIA). A series of 100 compds. were suggested. As di-Ph sulfide derivs. like [11C]DASB have recently proven to be promising PET ligands, rational modification of the di-Ph sulfide scaffold has been performed. The novel compds. were predicted to be selective high affinity SERT ligands. Important new ideas are the introduction of a fluoroethyl-oxycarbonyl group (ester) and fluoroethyl-carbonyl group (ketone), as well as a formyl group (aldehyde), and its corresponding oxime and imine. Another innovative suggestion is the replacement of the sulfur bridge with a cyanamide group and a fluoroethylamino group. The suggested compds. possess features providing new possibilities for carbon-11 or fluorine-18 labeling. Synthesis, biol. testing, and screening for PET suitability are reasonably further steps.

IT 622399-66-0 622399-67-1 622399-68-2

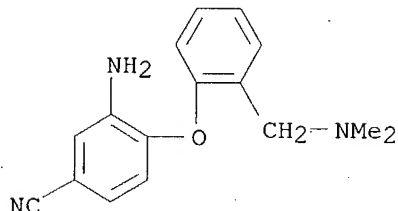
622399-69-3

RL: BSU (Biological study, unclassified); PRP (Properties); BIOL (Biological study)

(mol. modeling of PET radiotracers for serotonin transporter)

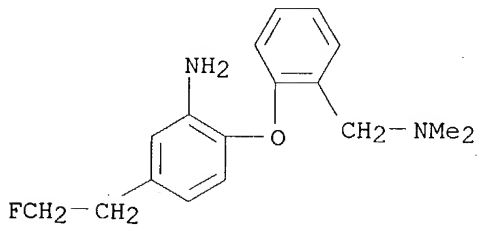
RN 622399-66-0 HCPLUS

CN Benzonitrile, 3-amino-4-[2-[(dimethylamino)methyl]phenoxy]- (9CI) (CA INDEX NAME)



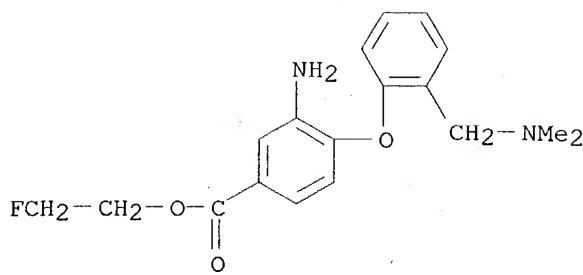
RN 622399-67-1 HCPLUS

CN Benzenemethanamine, 2-[2-amino-4-(2-fluoroethyl)phenoxy]-N,N-dimethyl- (9CI) (CA INDEX NAME)

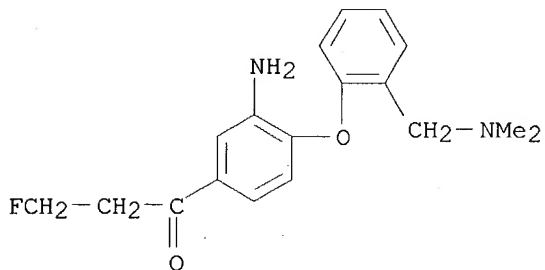


RN 622399-68-2 HCPLUS

CN Benzoic acid, 3-amino-4-[2-[(dimethylamino)methyl]phenoxy]-, 2-fluoroethyl ester (9CI) (CA INDEX NAME)



RN 622399-69-3 HCPLUS
 CN 1-Propanone, 1-[3-amino-4-[2-[(dimethylamino)methyl]phenoxy]phenyl]-3-fluoro- (9CI) (CA INDEX NAME)



REFERENCE COUNT: 59 THERE ARE 59 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L51 ANSWER 7 OF 44 HCPLUS COPYRIGHT 2004 ACS on STN
 ACCESSION NUMBER: 2002:925264 HCPLUS
 DOCUMENT NUMBER: 138:11431
 TITLE: 5-HT1a antagonist or an α 2-adrenergic antagonist in combination with an serotonin reuptake inhibitor for treatment of sleep disorders, including sleep apnea
 INVENTOR(S): Howard, Harry Ralph, Jr.
 PATENT ASSIGNEE(S): Pfizer Products Inc., USA
 SOURCE: Eur. Pat. Appl., 22 pp.
 CODEN: EPXXDW
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 1262197	A2	20021204	EP 2002-253589	20020522
EP 1262197	A3	20021218		
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR				

US 2002183306	A1	20021205	US 2002-75849	20020213
BR 2002001974	A	20030422	BR 2002-1974	20020528
JP 2003026602	A2	20030129	JP 2002-155222	20020529
			US 2001-294322P	P 20010530

PRIORITY APPLN. INFO.:

MARPAT 138:11431

AB The invention provides a method of treating sleep disorders, including sleep apnea, in a mammal, including a human, by administering to the mammal a 5-HT_{1a} antagonist or an α₂-adrenergic antagonist in combination with an serotonin reuptake inhibitor (SRI) antidepressant agent with improvement in efficacy. Also provided are pharmaceutical compns. containing a pharmaceutically acceptable carrier, a 5-HT_{1a} antagonist or an α₂-adrenergic antagonist, and an SRI antidepressant agent.

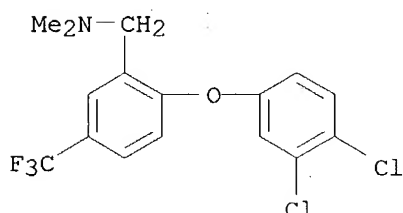
IT 289716-79-6 289716-94-5 289717-01-7
 289717-16-4 289717-18-6 289717-48-2
 289717-50-6 289717-52-8 289717-56-2
 289717-57-3 289717-59-5 289717-60-8
 289717-61-9 289717-62-0 289717-63-1
 289717-64-2 289717-65-3 289717-66-4
 289717-67-5 289717-68-6 289717-69-7
 289717-70-0 289717-71-1 289717-72-2
 289717-73-3 289717-74-4 289717-75-5
 444888-21-5 444888-22-6 444888-23-7
 444888-24-8 444888-25-9 444888-27-1
 444888-29-3 444888-31-7 444888-33-9
 444888-34-0 444888-35-1 444888-36-2
 444888-37-3 444888-38-4 444888-39-5
 444888-40-8 444888-41-9 444888-42-0
 444888-43-1 444888-45-3 444888-46-4
 444888-49-7 454456-66-7 477337-55-6

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(5-HT_{1a} antagonist or α₂-adrenergic antagonist in combination with serotonin reuptake inhibitor for treatment of sleep disorders, including sleep apnea)

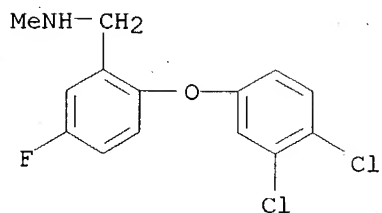
RN 289716-79-6 HCAPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-5-(trifluoromethyl)- (9CI) (CA INDEX NAME)

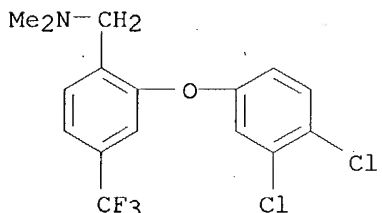


RN 289716-94-5 HCAPLUS

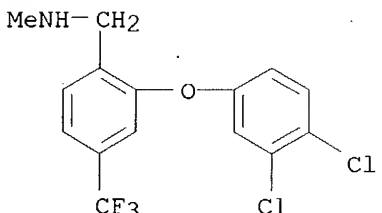
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N-methyl- (9CI) (CA INDEX NAME)



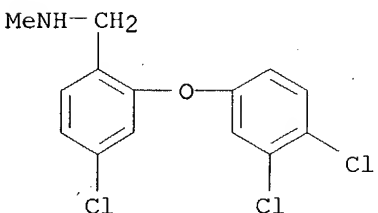
RN 289717-01-7 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-4-(trifluoromethyl)- (9CI) (CA INDEX NAME)



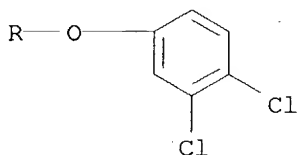
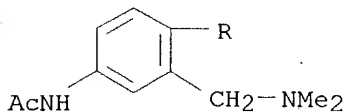
RN 289717-16-4 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-4-(trifluoromethyl)- (9CI) (CA INDEX NAME)



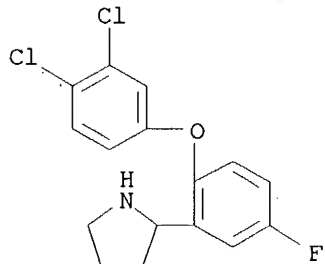
RN 289717-18-6 HCPLUS
 CN Benzenemethanamine, 4-chloro-2-(3,4-dichlorophenoxy)-N-methyl- (9CI) (CA INDEX NAME)



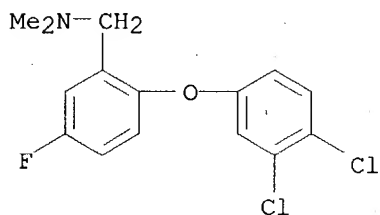
RN 289717-48-2 HCPLUS

CN Acetamide, N-[4-(3,4-dichlorophenoxy)-3-[(dimethylamino)methyl]phenyl]-
(9CI) (CA INDEX NAME)

RN 289717-50-6 HCPLUS

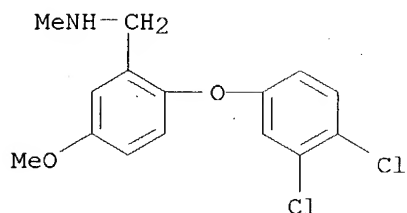
CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]- (9CI) (CA INDEX
NAME)

RN 289717-52-8 HCPLUS

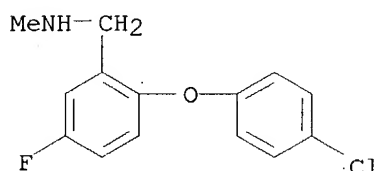
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N,N-dimethyl- (9CI)
(CA INDEX NAME)

RN 289717-56-2 HCPLUS

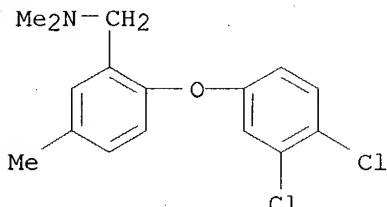
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-methoxy-N-methyl- (9CI) (CA
INDEX NAME)



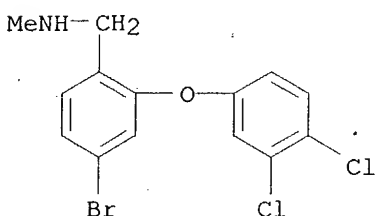
RN 289717-57-3 HCPLUS
 CN Benzenemethanamine, 2-(4-chlorophenoxy)-5-fluoro-N-methyl- (9CI) (CA
 INDEX NAME)



RN 289717-59-5 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N,5-trimethyl- (9CI) (CA
 INDEX NAME)

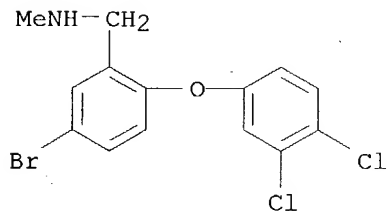


RN 289717-60-8 HCPLUS
 CN Benzenemethanamine, 4-bromo-2-(3,4-dichlorophenoxy)-N-methyl- (9CI) (CA
 INDEX NAME)



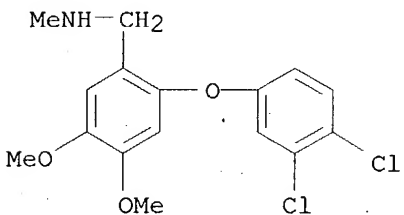
RN 289717-61-9 HCPLUS

CN Benzenemethanamine, 5-bromo-2-(3,4-dichlorophenoxy)-N-methyl- (9CI) (CA INDEX NAME)



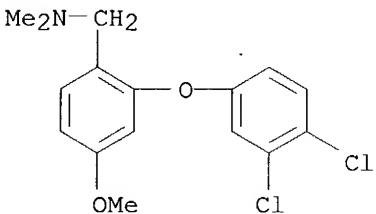
RN 289717-62-0 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4,5-dimethoxy-N-methyl- (9CI) (CA INDEX NAME)



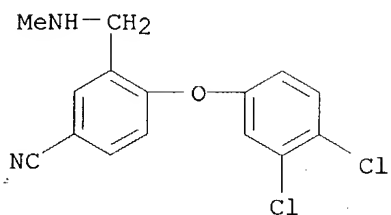
RN 289717-63-1 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4-methoxy-N,N-dimethyl- (9CI) (CA INDEX NAME)



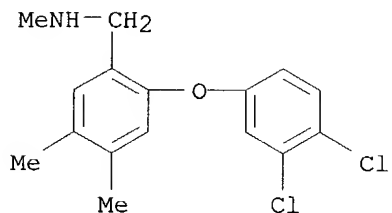
RN 289717-64-2 HCPLUS

CN Benzonitrile, 4-(3,4-dichlorophenoxy)-3-[(methylamino)methyl]- (9CI) (CA INDEX NAME)



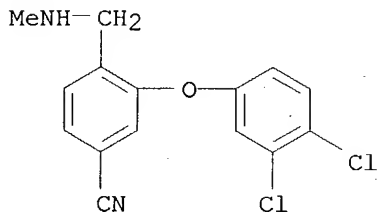
RN 289717-65-3 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,4,5-trimethyl- (9CI) (CA INDEX NAME)



RN 289717-66-4 HCPLUS

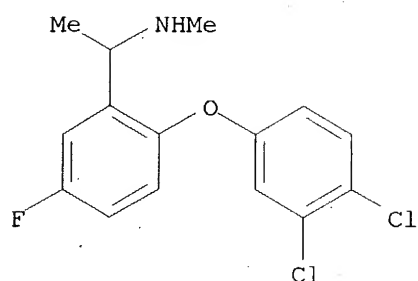
CN Benzonitrile, 3-(3,4-dichlorophenoxy)-4-[(methylamino)methyl]- (9CI) (CA INDEX NAME)



RN 289717-67-5 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N, α -dimethyl-, (+)- (9CI) (CA INDEX NAME)

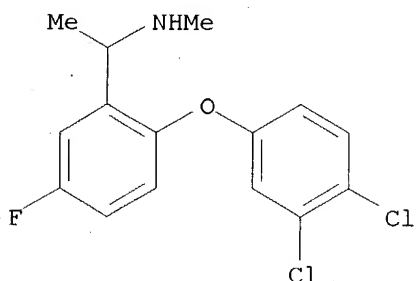
Rotation (+).



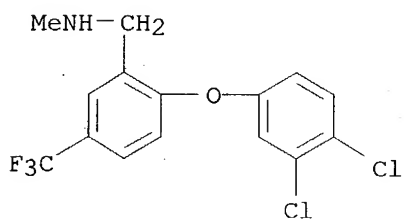
RN 289717-68-6 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N, α -dimethyl-,
(-) (9CI) (CA INDEX NAME)

Rotation (-).

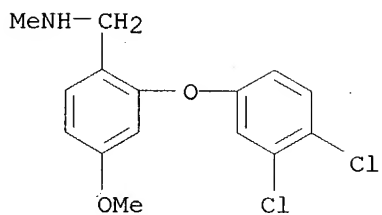


RN 289717-69-7 HCPLUS

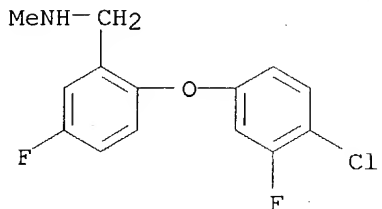
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-5-(trifluoromethyl)-
(9CI) (CA INDEX NAME)

RN 289717-70-0 HCPLUS

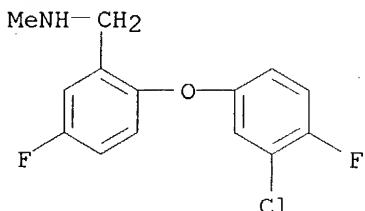
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4-methoxy-N-methyl- (9CI) (CA
INDEX NAME)



RN 289717-71-1 HCPLUS

CN Benzenemethanamine, 2-(4-chloro-3-fluorophenoxy)-5-fluoro-N-methyl- (9CI)
(CA INDEX NAME)

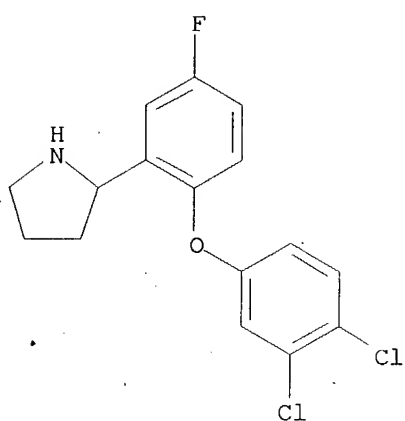
RN 289717-72-2 HCPLUS

CN Benzenemethanamine, 2-(3-chloro-4-fluorophenoxy)-5-fluoro-N-methyl- (9CI)
(CA INDEX NAME)

RN 289717-73-3 HCPLUS

CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-, (-)- (9CI) (CA INDEX NAME)

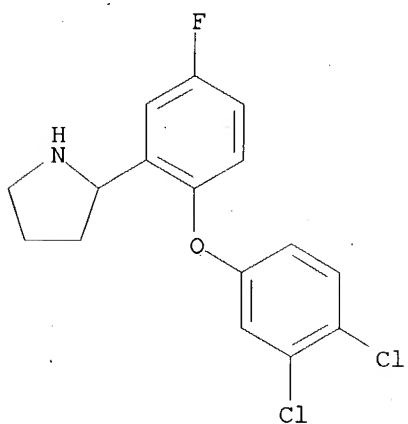
Rotation (-).



RN 289717-74-4 HCAPLUS

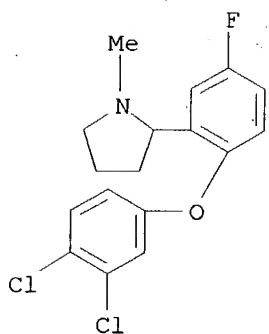
CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-, (+)- (9CI) (CA INDEX NAME)

Rotation (+).

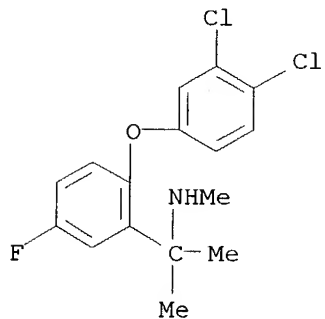


RN 289717-75-5 HCAPLUS

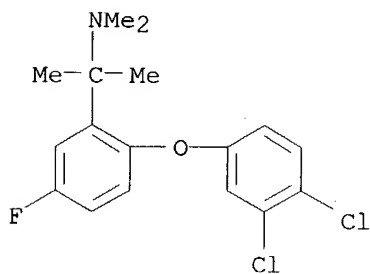
CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-1-methyl- (9CI) (CA INDEX NAME)



RN 444888-21-5 HCPLUS

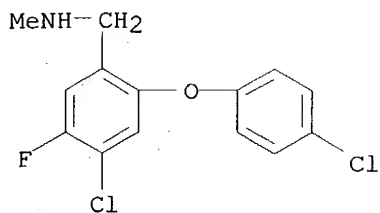
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N, α , α -trimethyl- (9CI) (CA INDEX NAME)

RN 444888-22-6 HCPLUS

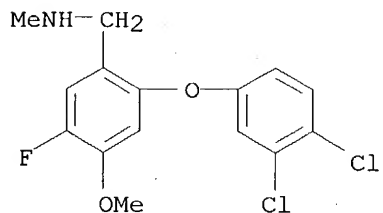
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N,N, α , α -tetramethyl- (9CI) (CA INDEX NAME)

RN 444888-23-7 HCPLUS

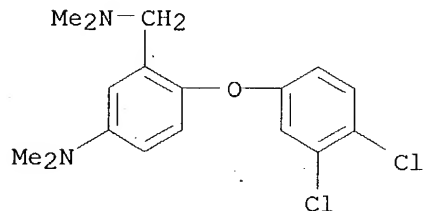
CN Benzenemethanamine, 4-chloro-2-(4-chlorophenoxy)-5-fluoro-N-methyl- (9CI) (CA INDEX NAME)



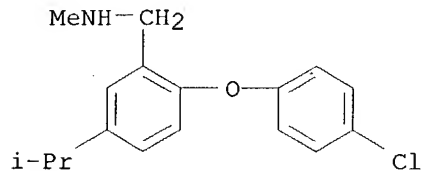
RN 444888-24-8 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-4-methoxy-N-methyl-
 (9CI) (CA INDEX NAME)



RN 444888-25-9 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-(dimethylamino)-N,N-dimethyl-
 (9CI) (CA INDEX NAME)

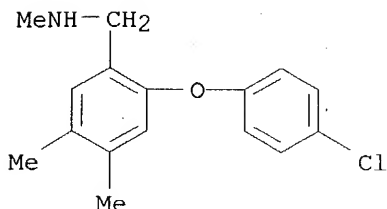


RN 444888-27-1 HCPLUS
 CN Benzenemethanamine, 2-(4-chlorophenoxy)-N-methyl-5-(1-methylethyl)- (9CI)
 (CA INDEX NAME)



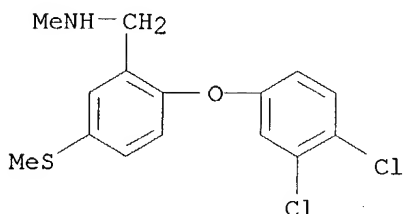
RN 444888-29-3 HCPLUS

CN Benzenemethanamine, 2-(4-chlorophenoxy)-N,4,5-trimethyl- (9CI) (CA INDEX NAME)

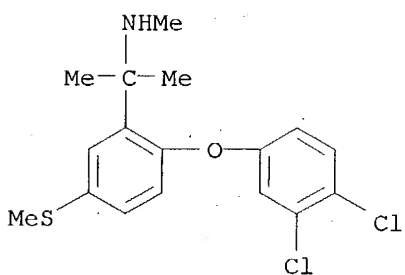


RN 444888-31-7 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-5-(methylthio)- (9CI) (CA INDEX NAME)

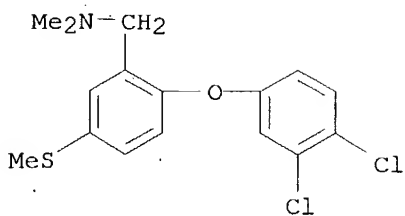


RN 444888-33-9 HCPLUS

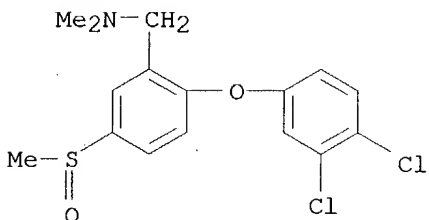
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N, α , α -trimethyl-5-(methylthio)- (9CI) (CA INDEX NAME)

RN 444888-34-0 HCPLUS

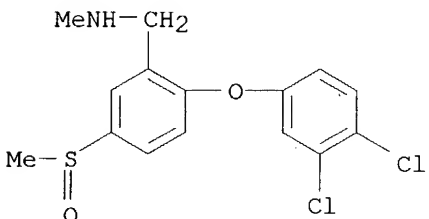
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-5-(methylthio)- (9CI) (CA INDEX NAME)



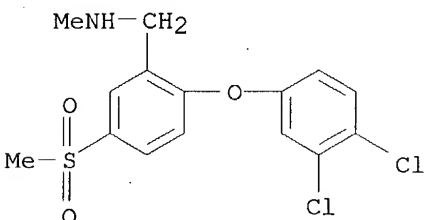
RN 444888-35-1 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-5-(methylsulfinyl)- (9CI) (CA INDEX NAME)



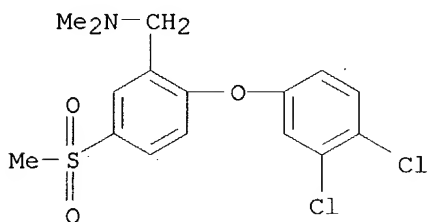
RN 444888-36-2 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-5-(methylsulfinyl)- (9CI) (CA INDEX NAME)



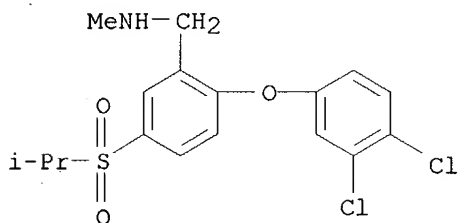
RN 444888-37-3 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-5-(methylsulfonyl)- (9CI) (CA INDEX NAME)



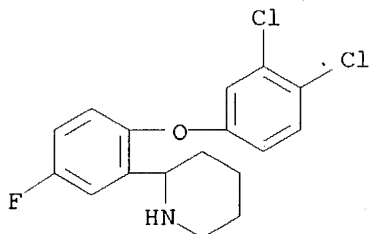
RN 444888-38-4 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-5-(methylsulfonyl)- (9CI) (CA INDEX NAME)



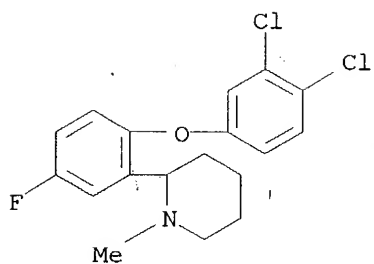
RN 444888-39-5 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-5-[(1-methylethyl)sulfonyl]- (9CI) (CA INDEX NAME)



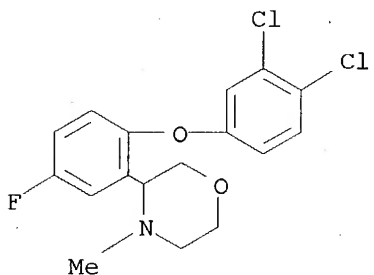
RN 444888-40-8 HCPLUS
 CN Piperidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]- (9CI) (CA INDEX NAME)



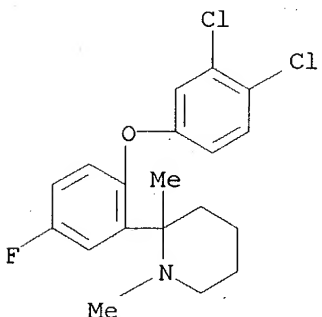
RN 444888-41-9 HCPLUS
 CN Piperidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-1-methyl- (9CI) (CA INDEX NAME)



RN 444888-42-0 HCPLUS

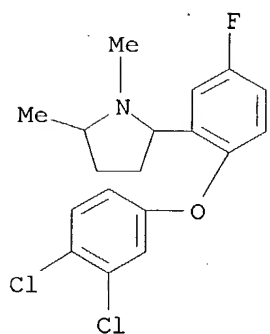
CN Morpholine, 3-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-4-methyl- (9CI)
(CA INDEX NAME)

RN 444888-43-1 HCPLUS

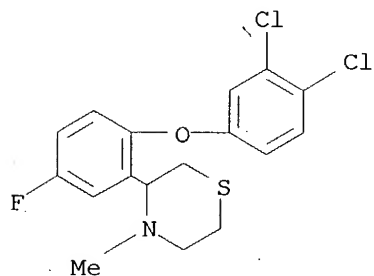
CN Piperidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-1,2-dimethyl- (9CI)
(CA INDEX NAME)

RN 444888-45-3 HCPLUS

CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-1,5-dimethyl-
(9CI) (CA INDEX NAME)

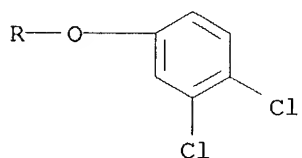
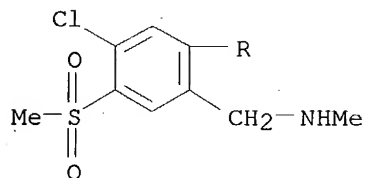


RN 444888-46-4 HCPLUS

CN Thiomorpholine, 3-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-4-methyl- (9CI)
(CA INDEX NAME)

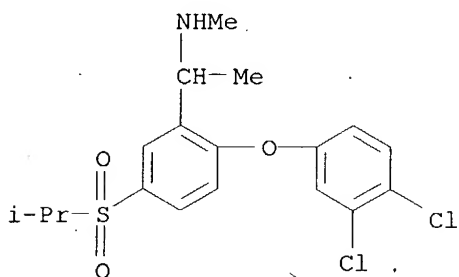
RN 444888-49-7 HCPLUS

CN Benzenemethanamine, 4-chloro-2-(3,4-dichlorophenoxy)-N-methyl-5-(methylsulfonyl)- (9CI) (CA INDEX NAME)



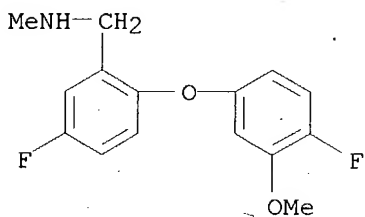
RN 454456-66-7 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N, α -dimethyl-5-[(1-methylethyl)sulfonyl]- (9CI) (CA INDEX NAME)



RN 477337-55-6 HCAPLUS

CN Benzenemethanamine, 5-fluoro-2-(4-fluoro-3-methoxyphenoxy)-N-methyl- (9CI)
(CA INDEX NAME)



L51 ANSWER 8 OF 44 HCAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: 2002:925263 HCAPLUS

DOCUMENT NUMBER: 138:336

TITLE: Combination of a monoamine reuptake inhibitor and an opioid antagonist for use in alcoholism and alcohol dependence

INVENTOR(S): Howard, Harry Ralph, Jr.

PATENT ASSIGNEE(S): Pfizer Products Inc., USA

SOURCE: Eur. Pat. Appl., 37 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 1262196	A2	20021204	EP 2002-253105	20020502
EP 1262196	A3	20021218		
		R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR		
JP 2002370975	A2	20021224	JP 2002-132804	20020508
AU 2002040686	A5	20021205	AU 2002-40686	20020516

US 2003130322
CN 1386503

A1 20030710
A 20021225

US 2002-153379
CN 2002-120350

20020522
20020523

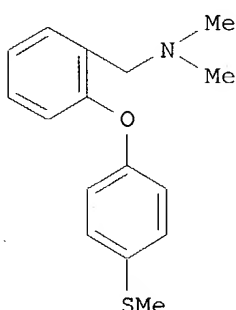
PRIORITY APPLN. INFO.:

US 2001-293088P P 20010523

OTHER SOURCE(S):

MARPAT 138:336

GI



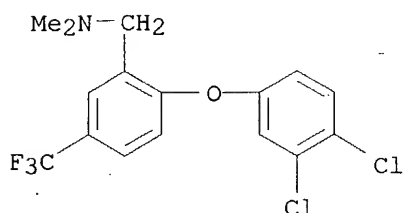
AB The present invention relates to a method of treating alcoholism or alc. dependence in a mammal, including a human, by administering to the mammal a monoamine reuptake inhibitor in combination with an opioid antagonist. It also relates to pharmaceutical compns. containing a pharmaceutically acceptable carrier, a monoamine reuptake inhibitor and an opioid antagonist. An example monoamine reuptake inhibitor is I.

IT 289716-79-6 289716-94-5 289717-01-7
289717-16-4 289717-18-6 289717-24-4
289717-48-2 289717-50-6 289717-52-8
289717-53-9 289717-54-0 289717-55-1
289717-56-2 289717-57-3 289717-58-4
289717-59-5 289717-60-8 289717-61-9
289717-62-0 289717-63-1 289717-64-2
289717-65-3 289717-66-4 289717-67-5
289717-68-6 289717-69-7 289717-70-0
289717-71-1 289717-72-2 289717-73-3
289717-74-4 289717-75-5 364323-82-0
476310-75-5

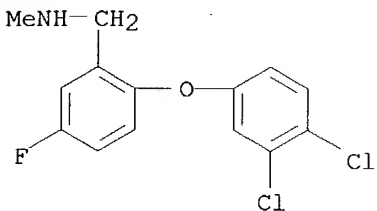
RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(combination of a monoamine reuptake inhibitor and an opioid antagonist
for use in alcoholism and alc. dependence)

RN 289716-79-6 HCAPLUS

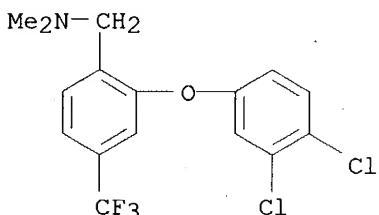
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-5-(trifluoromethyl)- (9CI) (CA INDEX NAME)



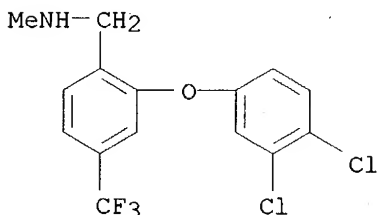
RN 289716-94-5 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N-methyl- (9CI) (CA INDEX NAME)



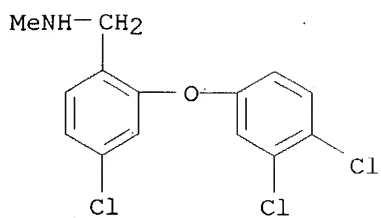
RN 289717-01-7 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-4-(trifluoromethyl)- (9CI) (CA INDEX NAME)



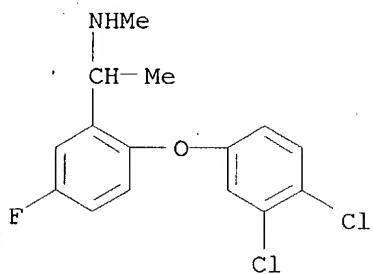
RN 289717-16-4 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-4-(trifluoromethyl)- (9CI) (CA INDEX NAME)



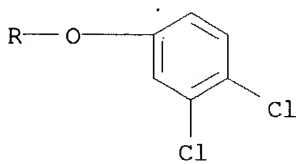
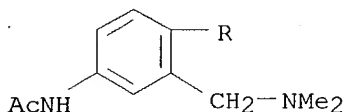
RN 289717-18-6 HCPLUS
 CN Benzenemethanamine, 4-chloro-2-(3,4-dichlorophenoxy)-N-methyl- (9CI) (CA INDEX NAME)



RN 289717-24-4 HCPLUS

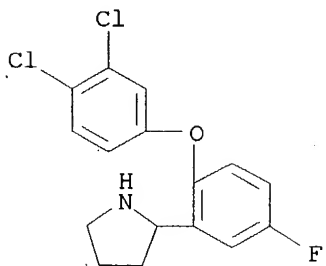
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N, α -dimethyl-
(9CI) (CA INDEX NAME)

RN 289717-48-2 HCPLUS

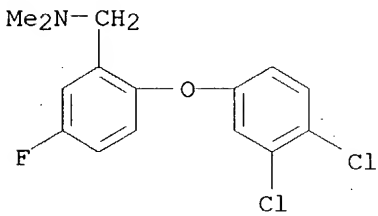
CN Acetamide, N-[4-(3,4-dichlorophenoxy)-3-(dimethylamino)methyl]phenyl-
(9CI) (CA INDEX NAME)

RN 289717-50-6 HCPLUS

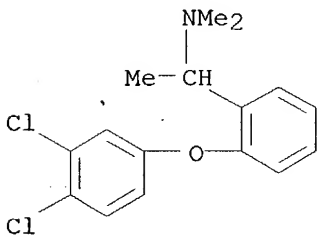
CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]- (9CI) (CA INDEX
NAME)



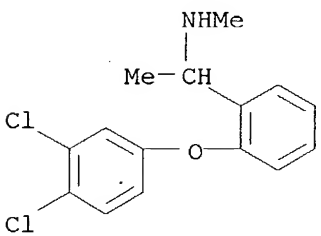
RN 289717-52-8 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N,N-dimethyl- (9CI)
(CA INDEX NAME)

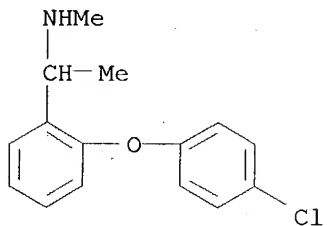
RN 289717-53-9 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N,α-trimethyl- (9CI)
(CA INDEX NAME)

RN 289717-54-0 HCPLUS

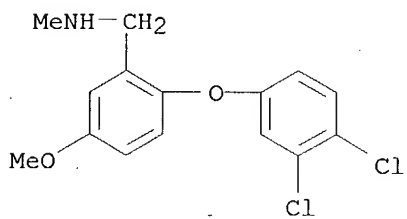
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,α-dimethyl- (9CI) (CA
INDEX NAME)

RN 289717-55-1 HCAPLUS

CN Benzenemethanamine, 2-(4-chlorophenoxy)-N, α -dimethyl- (9CI) (CA INDEX NAME)

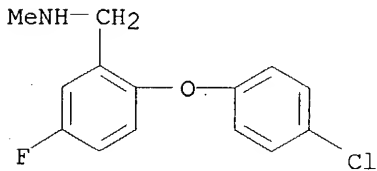
RN 289717-56-2 HCAPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-methoxy-N-methyl- (9CI) (CA INDEX NAME)



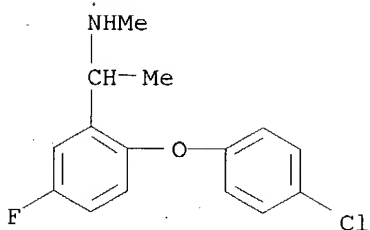
RN 289717-57-3 HCAPLUS

CN Benzenemethanamine, 2-(4-chlorophenoxy)-5-fluoro-N-methyl- (9CI) (CA INDEX NAME)



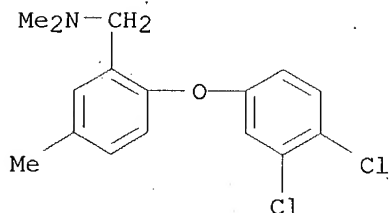
RN 289717-58-4 HCAPLUS

CN Benzenemethanamine, 2-(4-chlorophenoxy)-5-fluoro-N, α -dimethyl- (9CI) (CA INDEX NAME)



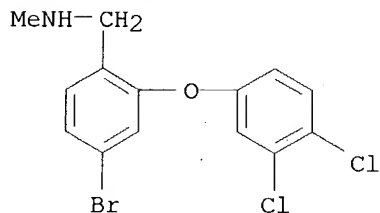
RN 289717-59-5 HCAPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N,5-trimethyl- (9CI) (CA INDEX NAME)



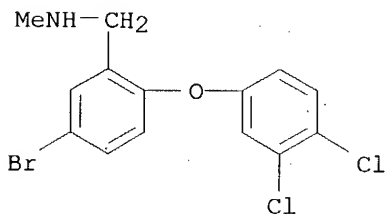
RN 289717-60-8 HCAPLUS

CN Benzenemethanamine, 4-bromo-2-(3,4-dichlorophenoxy)-N-methyl- (9CI) (CA INDEX NAME)

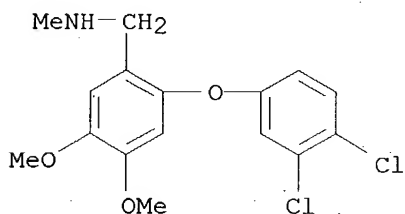


RN 289717-61-9 HCAPLUS

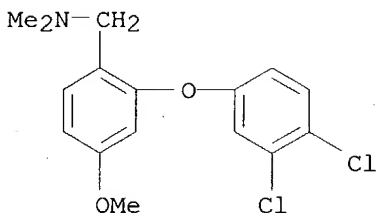
CN Benzenemethanamine, 5-bromo-2-(3,4-dichlorophenoxy)-N-methyl- (9CI) (CA INDEX NAME)



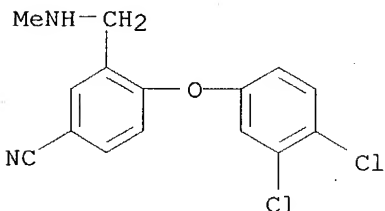
RN 289717-62-0 HCAPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4,5-dimethoxy-N-methyl- (9CI)
(CA INDEX NAME)

RN 289717-63-1 HCAPLUS

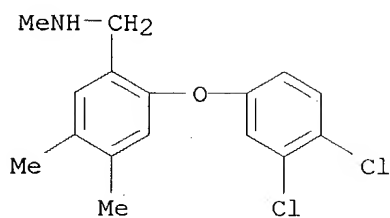
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4-methoxy-N,N-dimethyl- (9CI)
(CA INDEX NAME)

RN 289717-64-2 HCAPLUS

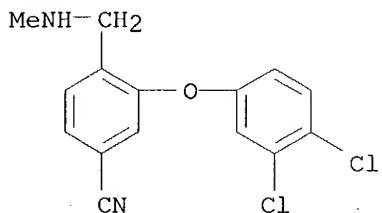
CN Benzonitrile, 4-(3,4-dichlorophenoxy)-3-[(methylamino)methyl]- (9CI) (CA
INDEX NAME)

RN 289717-65-3 HCAPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,4,5-trimethyl- (9CI) (CA
INDEX NAME)

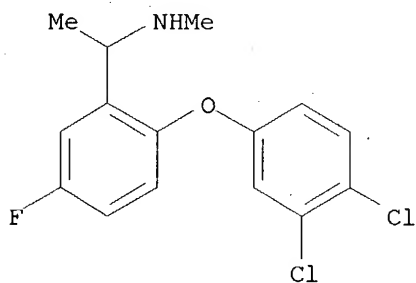


RN 289717-66-4 HCAPLUS
 CN Benzonitrile, 3-(3,4-dichlorophenoxy)-4-[(methylamino)methyl]- (9CI) (CA INDEX NAME)



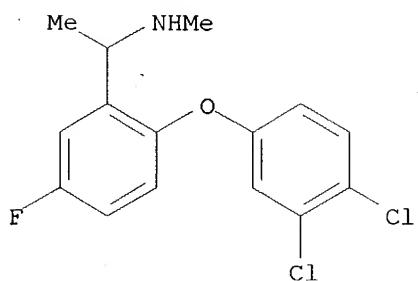
RN 289717-67-5 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N, α -dimethyl-, (+)- (9CI) (CA INDEX NAME)

Rotation (+).

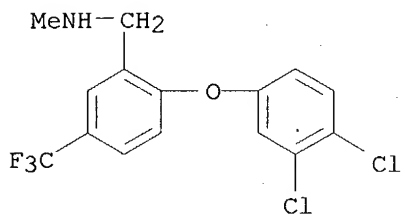


RN 289717-68-6 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N, α -dimethyl-, (-)- (9CI) (CA INDEX NAME)

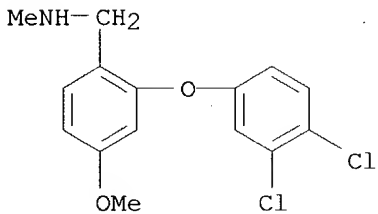
Rotation (-).



RN 289717-69-7 HCAPLUS

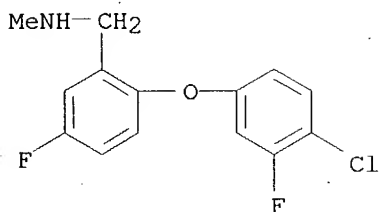
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-5-(trifluoromethyl)-
(9CI) (CA INDEX NAME)

RN 289717-70-0 HCAPLUS

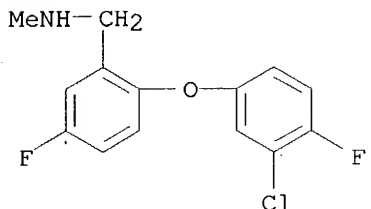
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4-methoxy-N-methyl- (9CI) (CA
INDEX NAME)

RN 289717-71-1 HCAPLUS

CN Benzenemethanamine, 2-(4-chloro-3-fluorophenoxy)-5-fluoro-N-methyl- (9CI)
(CA INDEX NAME)



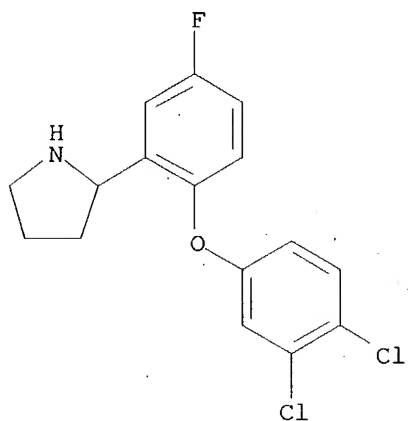
RN 289717-72-2 HCAPLUS

CN Benzenemethanamine, 2-(3-chloro-4-fluorophenoxy)-5-fluoro-N-methyl- (9CI)
(CA INDEX NAME)

RN 289717-73-3 HCAPLUS

CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-, (-)- (9CI) (CA
INDEX NAME)

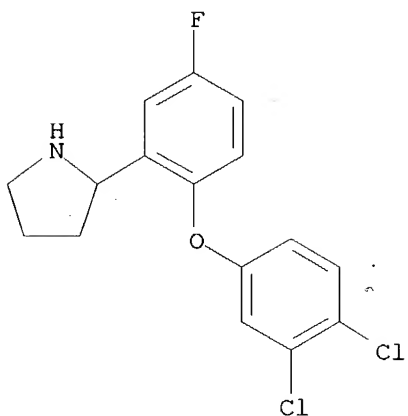
Rotation (-).



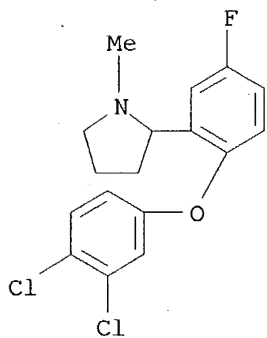
RN 289717-74-4 HCAPLUS

CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-, (+)- (9CI) (CA
INDEX NAME)

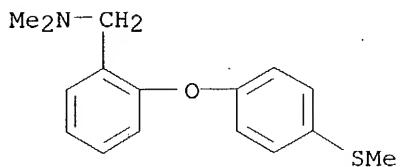
Rotation (+).



RN 289717-75-5 HCAPLUS

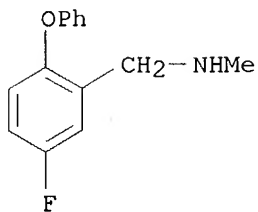
CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-1-methyl- (9CI)
(CA INDEX NAME)

RN 364323-82-0 HCAPLUS

CN Benzenemethanamine, N,N-dimethyl-2-[4-(methylthio)phenoxy]- (9CI) (CA
INDEX NAME)

RN 476310-75-5 HCAPLUS

CN Benzenemethanamine, 5-fluoro-N-methyl-2-phenoxy- (9CI) (CA INDEX NAME)



L51 ANSWER 9 OF 44 HCAPLUS COPYRIGHT 2004 ACS on STN
 ACCESSION NUMBER: 2002:904325 HCAPLUS
 DOCUMENT NUMBER: 137:380038
 TITLE: Combination treatment with monoamine reuptake inhibitor and dopamine D3 receptor agonist for depression and anxiety
 INVENTOR(S): Howard, Harry Ralph, Jr.
 PATENT ASSIGNEE(S): Pfizer Products Inc., USA
 SOURCE: Eur. Pat. Appl., 31 pp.
 CODEN: EPXXDW
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 1260221	A2	20021127	EP 2002-253135	20020503
EP 1260221	A3	20021218		
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR				
AU 2002040681	A5	20021205	AU 2002-40681	20020516
JP 2002370976	A2	20021224	JP 2002-141515	20020516
CN 1386504	A	20021225	CN 2002-120351	20020523

PRIORITY APPLN. INFO.: US 2001-293063P P 20010523

OTHER SOURCE(S): MARPAT 137:380038

AB The present invention relates to a method of treating depression or anxiety in a mammal, including a human, by administering to the mammal a monoamine reuptake inhibitor in combination with a dopamine D3 receptor agonist. It also relates to pharmaceutical compns. containing a pharmaceutically acceptable carrier, a monoamine reuptake inhibitor and a dopamine D3 receptor agonist.

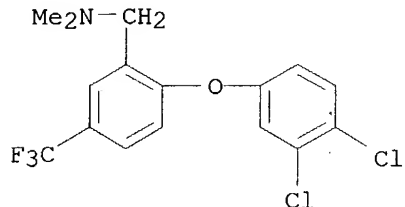
IT 289716-79-6 289716-94-5 289717-01-7
 289717-16-4 289717-18-6 289717-24-4
 289717-48-2 289717-50-6 289717-52-8
 289717-53-9 289717-54-0 289717-55-1
 289717-56-2 289717-57-3 289717-58-4
 289717-59-5 289717-60-8 289717-61-9
 289717-62-0 289717-63-1 289717-64-2
 289717-65-3 289717-66-4 289717-67-5
 289717-68-6 289717-69-7 289717-70-0
 289717-71-1 289717-72-2 289717-73-3
 289717-74-4 289717-75-5

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(monoamine reuptake inhibitor; combination treatment with monoamine reuptake inhibitor and dopamine D3 receptor agonist for depression and anxiety)

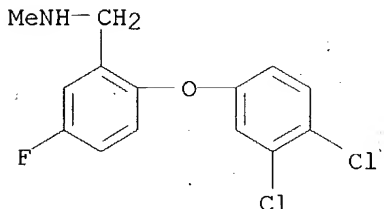
RN 289716-79-6 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-5-(trifluoromethyl)- (9CI) (CA INDEX NAME)



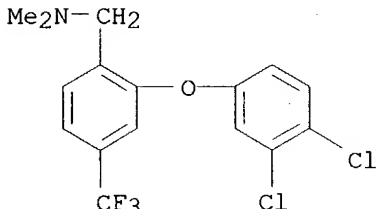
RN 289716-94-5 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N-methyl- (9CI) (CA INDEX NAME)



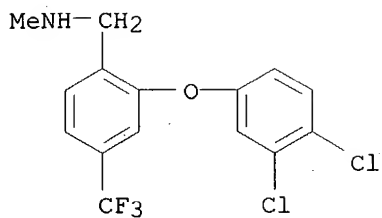
RN 289717-01-7 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-4-(trifluoromethyl)- (9CI) (CA INDEX NAME)

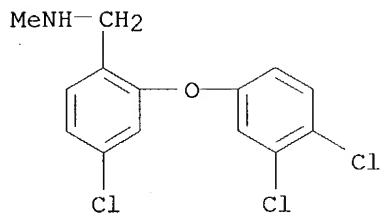


RN 289717-16-4 HCPLUS

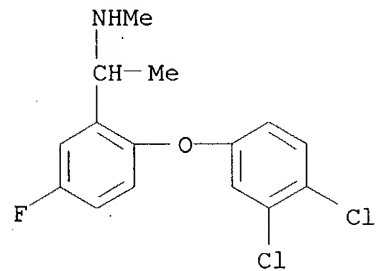
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-4-(trifluoromethyl)- (9CI) (CA INDEX NAME)



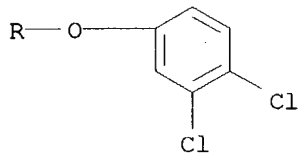
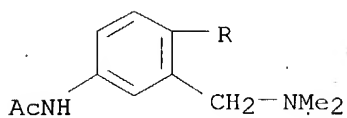
RN 289717-18-6 HCAPLUS
 CN Benzenemethanamine, 4-chloro-2-(3,4-dichlorophenoxy)-N-methyl- (9CI) (CA INDEX NAME)



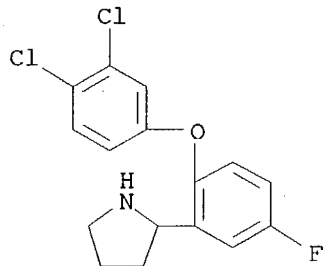
RN 289717-24-4 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N, α -dimethyl- (9CI) (CA INDEX NAME)



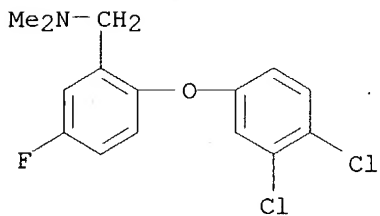
RN 289717-48-2 HCAPLUS
 CN Acetamide, N-[4-(3,4-dichlorophenoxy)-3-[(dimethylamino)methyl]phenyl]- (9CI) (CA INDEX NAME)



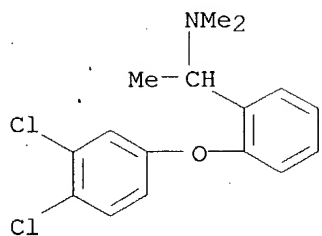
RN 289717-50-6 HCAPLUS
 CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]- (9CI) (CA INDEX NAME)



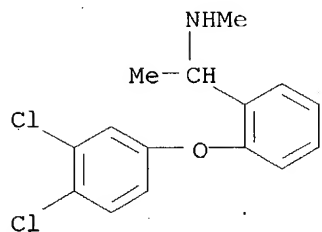
RN 289717-52-8 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N,N-dimethyl- (9CI)
 (CA INDEX NAME)



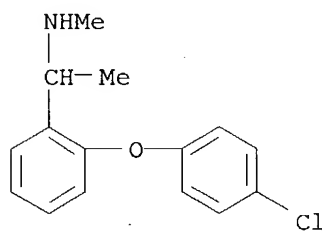
RN 289717-53-9 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N, α -trimethyl- (9CI)
 (CA INDEX NAME)



RN 289717-54-0 HCPLUS

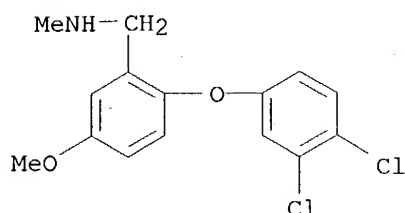
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N, α -dimethyl- (9CI) (CA INDEX NAME)

RN 289717-55-1 HCPLUS

CN Benzenemethanamine, 2-(4-chlorophenoxy)-N, α -dimethyl- (9CI) (CA INDEX NAME)

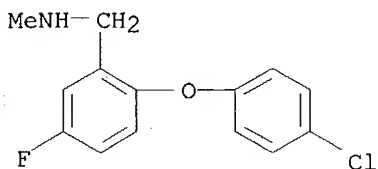
RN 289717-56-2 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-methoxy-N-methyl- (9CI) (CA INDEX NAME)

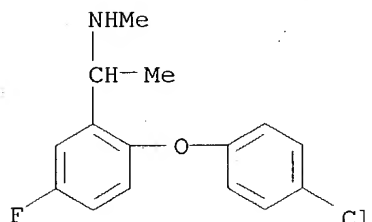


RN 289717-57-3 HCAPLUS

CN Benzenemethanamine, 2-(4-chlorophenoxy)-5-fluoro-N-methyl- (9CI) (CA INDEX NAME)

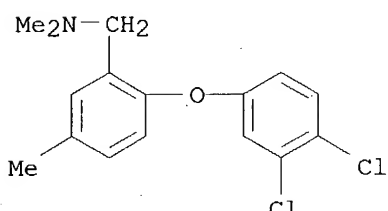


RN 289717-58-4 HCAPLUS

CN Benzenemethanamine, 2-(4-chlorophenoxy)-5-fluoro-N, α -dimethyl- (9CI) (CA INDEX NAME)

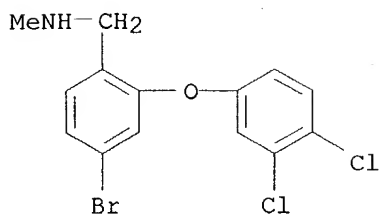
RN 289717-59-5 HCAPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N,5-trimethyl- (9CI) (CA INDEX NAME)

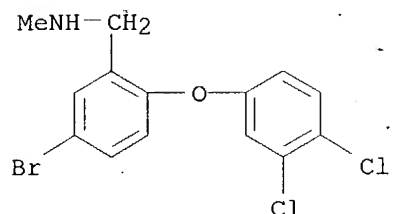


RN 289717-60-8 HCAPLUS

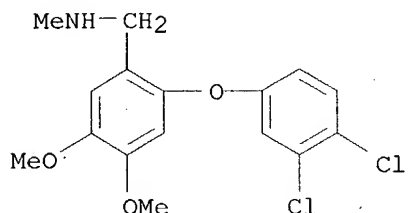
CN Benzenemethanamine, 4-bromo-2-(3,4-dichlorophenoxy)-N-methyl- (9CI) (CA INDEX NAME)



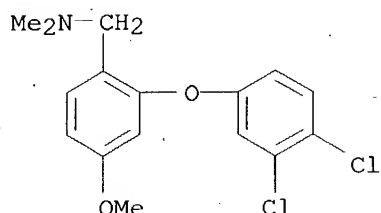
RN 289717-61-9 HCAPLUS
 CN Benzenemethanamine, 5-bromo-2-(3,4-dichlorophenoxy)-N-methyl- (9CI) (CA INDEX NAME)



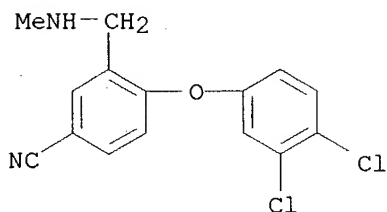
RN 289717-62-0 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4,5-dimethoxy-N-methyl- (9CI) (CA INDEX NAME)



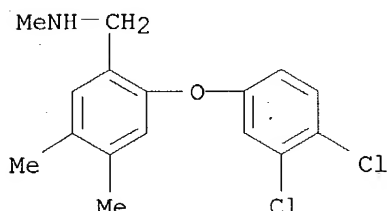
RN 289717-63-1 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4-methoxy-N,N-dimethyl- (9CI) (CA INDEX NAME)



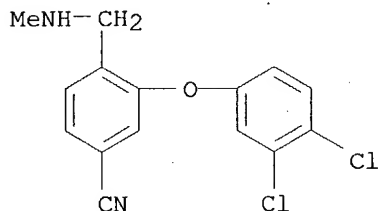
RN 289717-64-2 HCAPLUS
 CN Benzonitrile, 4-(3,4-dichlorophenoxy)-3-[(methylamino)methyl]- (9CI) (CA INDEX NAME)



RN 289717-65-3 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,4,5-trimethyl- (9CI) (CA INDEX NAME)

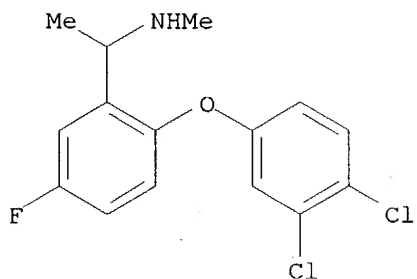


RN 289717-66-4 HCAPLUS
 CN Benzonitrile, 3-(3,4-dichlorophenoxy)-4-[(methylamino)methyl]- (9CI) (CA INDEX NAME)



RN 289717-67-5 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N, α -dimethyl-,
 (+)- (9CI) (CA INDEX NAME)

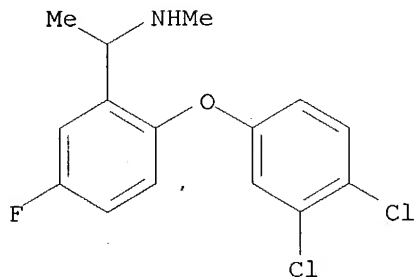
Rotation (+).



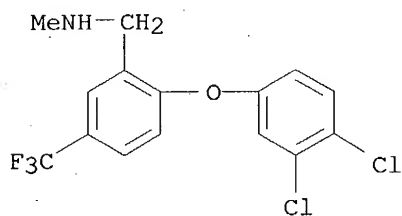
RN 289717-68-6 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-*N*,*α*-dimethyl-,
(-) (9CI) (CA INDEX NAME)

Rotation (-).

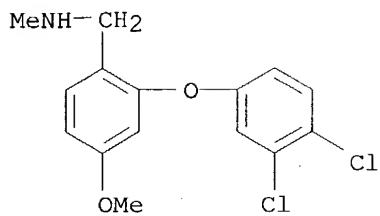


RN 289717-69-7 HCPLUS

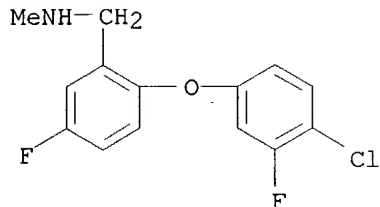
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-*N*-methyl-5-(trifluoromethyl)-
(9CI) (CA INDEX NAME)

RN 289717-70-0 HCPLUS

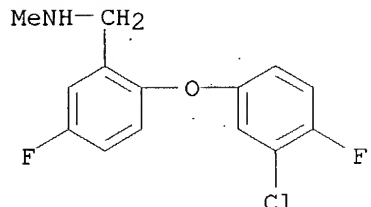
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4-methoxy-*N*-methyl- (9CI) (CA
INDEX NAME)



RN 289717-71-1 HCAPLUS
 CN Benzenemethanamine, 2-(4-chloro-3-fluorophenoxy)-5-fluoro-N-methyl- (9CI)
 (CA INDEX NAME)

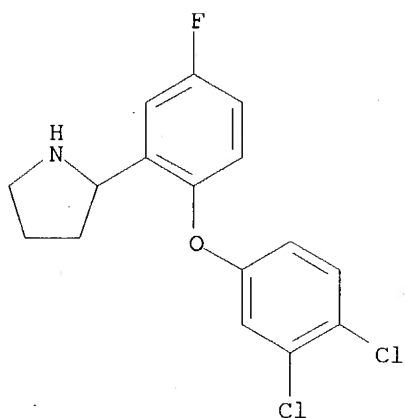


RN 289717-72-2 HCAPLUS
 CN Benzenemethanamine, 2-(3-chloro-4-fluorophenoxy)-5-fluoro-N-methyl- (9CI)
 (CA INDEX NAME)



RN 289717-73-3 HCAPLUS
 CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-, (-)- (9CI) (CA
 INDEX NAME)

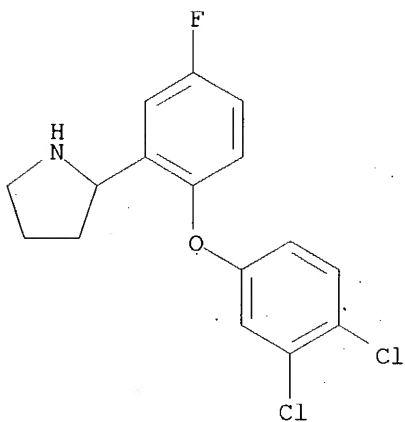
Rotation (-).



RN 289717-74-4 HCAPLUS

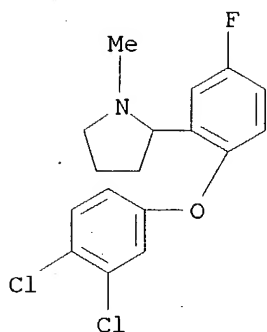
CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-, (+)- (9CI) (CA INDEX NAME)

Rotation (+).



RN 289717-75-5 HCAPLUS

CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-1-methyl- (9CI) (CA INDEX NAME)



L51 ANSWER 10 OF 44 HCAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: 2002:773622 HCAPLUS

DOCUMENT NUMBER: 137:273225

TITLE: Combination treatment for multiple sclerosis, other demyelinating conditions and peripheral neuropathy, especially painful neuropathies and diabetic neuropathy

INVENTOR(S): Howard, Harry Ralph, Jr.

PATENT ASSIGNEE(S): Pfizer Products Inc., USA

SOURCE: Eur. Pat. Appl., 15 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 1247533	A2	20021009	EP 2002-251844	20020314
EP 1247533	A3	20031217		
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR				
US 2002147206	A1	20021010	US 2001-24968	20011219
JP 2002356445	A2	20021213	JP 2002-100682	20020403
BR 2002001094	A	20030527	BR 2002-1094	20020405

PRIORITY APPLN. INFO.: US 2001-281988P P 20010405

OTHER SOURCE(S): MARPAT 137:273225

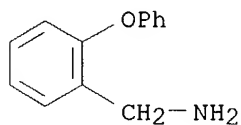
AB The invention relates to a combination useful in treating Multiple Sclerosis, other demyelinating disorders and peripheral neuropathy in a mammal comprising a neurotransmitter-inducing or precursor agent in combination with an (serotonin reuptake inhibitors, SRI) anxiolytic agent or an antidepressant with improvement in efficiency. It also relates to pharmaceutical compns. containing a pharmaceutically acceptable carrier, a neurotransmitter-inducing or precursor agent, and an SRI antidepressant or anxiolytic agent.

IT 107624-14-6 107624-14-6D, derivs.

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(combination treatment for multiple sclerosis, demyelinating conditions and peripheral neuropathy)

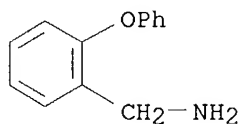
RN 107624-14-6 HCAPLUS

CN Benzenemethanamine, 2-phenoxy- (9CI) (CA INDEX NAME)



RN 107624-14-6 HCPLUS

CN Benzenemethanamine, 2-phenoxy- (9CI) (CA INDEX NAME)



L51 ANSWER 11 OF 44 HCPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: 2002:755211 HCPLUS

DOCUMENT NUMBER: 137:262839

TITLE: Preparation of phenoxybenzylamines as monoamine reuptake inhibitors for treatment of CNS disorders.

INVENTOR(S): Howard, Harry R.; Schmidt, Christopher J.; Seeger, Thomas F.; Elliott, Mark L.

PATENT ASSIGNEE(S): Pfizer, Inc., USA

SOURCE: U.S. Pat. Appl. Publ., 24 pp., Cont.-in-part of U.S. Ser. No. 529,207.

CODEN: USXXCO

DOCUMENT TYPE: Patent

LANGUAGE: English

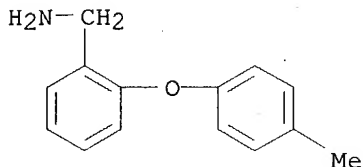
FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2002143003	A1	20021003	US 2001-845992	20010430
US 6677378	B2	20040113		
WO 2000050380	A1	20000831	WO 2000-IB108	20000202
W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM				
RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG				
PRIORITY APPLN. INFO.:			US 1999-121313P	P 19990223
			US 2000-529207	A2 20000202
			WO 2000-IB108	W 20000202
OTHER SOURCE(S):	MARPAT	137:262839		
AB ROZCR3R4NR1R2 [R = (un)substituted Ph; R1,R2 = H, alk(en)yl, alkynyl;				

NR₁R₂ = heterocyclyl; R₃,R₄ = H or (fluoro)alkyl; R₃R₄ = (un)substituted alkylene; R₂R₃ = atoms to complete a heterocyclic ring; Z = (un)substituted phenylene] were prepared as monoamine reuptake inhibitors (no data). Such compds. are useful exhibit activity as serotonin, norepinephrine and dopamine reuptake inhibitors, and their pharmaceutically acceptable salts, and their use in the treatment of central nervous system and other disorders.

- IT 289718-11-2P, Benzenemethanamine, 2-(4-methylphenoxy)-
 RL: BYP (Byproduct); PREP (Preparation)
 (preparation of phenoxybenzylamines as monoamine reuptake inhibitors)
- RN 289718-11-2 HCPLUS
- CN Benzenemethanamine, 2-(4-methylphenoxy)- (9CI) (CA INDEX NAME)

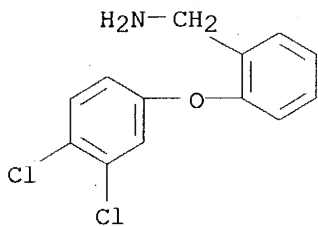


- IT 146520-69-6P, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-, hydrochloride 146797-20-8P, Benzenemethanamine, 2-(4-chlorophenoxy)-N,N-dimethyl-, (2Z)-2-butenedioate (1:1)
 289716-74-1P, Benzenemethanamine, 2-fluoro-6-(4-methylphenoxy)-
 289716-75-2P, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-
 289716-79-6P, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-5-(trifluoromethyl)- 289716-80-9P, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-5-(trifluoromethyl)-, (2Z)-2-butenedioate (1:1) 289716-82-1P, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-, hydrochloride 289716-85-4P, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-, (2Z)-2-butenedioate (4:3) 289716-88-7P, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-ethyl-, (2Z)-2-butenedioate (1:1)
 289716-89-8P, Benzenemethanamine, 2-(4-chlorophenoxy)-5-fluoro-N-methyl-, hydrochloride 289716-91-2P, Benzenemethanamine, 2-(4-chlorophenoxy)-N-methyl-, (2Z)-2-butenedioate (1:1)
 289716-92-3P, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N,N-dimethyl-, hydrochloride 289716-93-4P, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N-methyl-, hydrochloride
 289716-94-5P, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N-methyl- 289716-95-6P, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N-methyl-, (2Z)-2-butenedioate (2:1)
 289716-96-7P, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-5-nitro- 289716-97-8P, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-3-(trifluoromethyl)-, hydrochloride 289716-98-9P, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N, α -trimethyl-, hydrochloride 289717-00-6P, Benzenemethanamine, N,N-dimethyl-2-(4-methylphenoxy)-, (2Z)-2-butenedioate (1:1) 289717-01-7P, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-4-(trifluoromethyl)- 289717-02-8P, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-4-(trifluoromethyl)-, (2Z)-2-butenedioate (1:1) 289717-04-0P, Benzenemethanamine, 2-(3,4-difluorophenoxy)-N,N-dimethyl-, (2Z)-2-butenedioate (1:1) 289717-06-2P, Benzenemethanamine,

2-(3,4-difluorophenoxy)-N-methyl-, (2Z)-2-butenedioate (1:1)
289717-08-4P, Benzenemethanamine, N, α -dimethyl-2-(4-methylphenoxy)-, (2Z)-2-butenedioate (1:1) **289717-09-5P**, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-6-(trifluoromethyl)-, hydrochloride **289717-11-9P**, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-6-(trifluoromethyl)-, (2Z)-2-butenedioate (1:1) **289717-13-1P**, Benzenemethanamine, 2-(3,4-dimethylphenoxy)-5-fluoro-N-methyl-, (2Z)-2-butenedioate (1:1) **289717-15-3P**, Benzenemethanamine, 2-(3,4-dimethylphenoxy)-N-methyl-, (2Z)-2-butenedioate (1:1) **289717-16-4P**, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-4-(trifluoromethyl)- **289717-17-5P**, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-4-(trifluoromethyl)-, (2Z)-2-butenedioate (1:1) **289717-18-6P**, Benzenemethanamine, 4-chloro-2-(3,4-dichlorophenoxy)-N-methyl- **289717-19-7P**, Benzenemethanamine, 4-chloro-2-(3,4-dichlorophenoxy)-N-methyl-, (2Z)-2-butenedioate (1:1) **289717-23-3P**, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-3-fluoro-N-methyl-, (2Z)-2-butenedioate (1:1) **289717-24-4P**, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N, α -dimethyl- **289717-25-5P**, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N, α -dimethyl-, (2Z)-2-butenedioate (1:1) **289717-26-6P**, Benzenemethanamine, 2-(4-chlorophenoxy)-5-fluoro-N, α -dimethyl-, hydrochloride **289717-28-8P**, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4-fluoro-N-methyl-, (2Z)-2-butenedioate (1:1) **289717-29-9P**, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-methoxy-N-methyl-, hydrochloride **289717-30-2P**, Benzenemethanamine, 2-(4-chlorophenoxy)-N, α -dimethyl-, hydrochloride **289717-32-4P**, Benzenemethanamine, 2-(4-chlorophenoxy)-5-fluoro-N,N-dimethyl-, (2Z)-2-butenedioate (1:1) **289717-33-5P**, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4,5-dimethoxy-N-methyl-, hydrochloride **289717-34-6P**, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,4,5-trimethyl-, hydrochloride **289717-35-7P**, Benzenemethanamine, 5-bromo-2-(3,4-dichlorophenoxy)-N-methyl-, hydrochloride **289717-36-8P**, Benzenemethanamine, 4-bromo-2-(3,4-dichlorophenoxy)-N-methyl-, hydrochloride **289717-37-9P**, Benzenemethanamine, N,N-dimethyl-2-[4-(methylthio)phenoxy]-, hydrochloride **289717-38-0P**, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-, hydrochloride **289717-39-1P**, Benzonitrile, 3-(3,4-dichlorophenoxy)-4-[(methylamino)methyl]-, monohydrochloride **289717-41-5P**, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4-methoxy-N,N-dimethyl-, hydrochloride **289717-42-6P**, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N,5-trimethyl-, hydrochloride **289717-43-7P**, Benzenemethanamine, 2-fluoro-N,N-dimethyl-6-(4-methylphenoxy)- **289717-44-8P**, Benzenemethanamine, 2-fluoro-N,N-dimethyl-6-(4-methylphenoxy)-, (2Z)-2-butenedioate (1:1) **289717-45-9P**, Benzenemethanamine, 5-amino-2-(3,4-dichlorophenoxy)-N,N-dimethyl-, monohydrochloride **289717-46-0P**, Benzenemethanamine, 5-amino-2-(3,4-dichlorophenoxy)-N,N-dimethyl- **289717-47-1P**, Acetamide, N-[4-(3,4-dichlorophenoxy)-3-[(dimethylamino)methyl]phenyl]-, monohydrochloride **289717-48-2P**, Acetamide, N-[4-(3,4-dichlorophenoxy)-3-[(dimethylamino)methyl]phenyl]- **289717-49-3P**, Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-, hydrochloride **289717-50-6P**, Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]- **289717-51-7P**, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-, hydrochloride **289717-52-8P**, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N,N-dimethyl-

289717-53-9P, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N, α -trimethyl- **289717-54-0P**, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N, α -dimethyl- **289717-55-1P**, Benzenemethanamine, 2-(4-chlorophenoxy)-N, α -dimethyl-
289717-56-2P, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-methoxy-N-methyl- **289717-57-3P**, Benzenemethanamine, 2-(4-chlorophenoxy)-5-fluoro-N-methyl- **289717-58-4P**, Benzenemethanamine, 2-(4-chlorophenoxy)-5-fluoro-N, α -dimethyl-
289717-59-5P, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N,5-trimethyl- **289717-60-8P**, Benzenemethanamine, 4-bromo-2-(3,4-dichlorophenoxy)-N-methyl- **289717-61-9P**, Benzenemethanamine, 5-bromo-2-(3,4-dichlorophenoxy)-N-methyl-
289717-62-0P, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4,5-dimethoxy-N-methyl- **289717-63-1P**, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4-methoxy-N,N-dimethyl- **289717-64-2P**, Benzonitrile, 4-(3,4-dichlorophenoxy)-3-[(methylamino)methyl]-
289717-65-3P, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,4,5-trimethyl- **289717-66-4P**, Benzonitrile, 3-(3,4-dichlorophenoxy)-4-[(methylamino)methyl]- **289717-67-5P**, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N, α -dimethyl-, (+)-
289717-68-6P, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N, α -dimethyl-, (-)- **289717-69-7P**, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-5-(trifluoromethyl)- **289717-70-0P**, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4-methoxy-N-methyl-
289717-71-1P, Benzenemethanamine, 2-(4-chloro-3-fluorophenoxy)-5-fluoro-N-methyl- **289717-72-2P**, Benzenemethanamine, 2-(3-chloro-4-fluorophenoxy)-5-fluoro-N-methyl- **289717-73-3P**, Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-, (-)-
289717-74-4P, Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-, (+)- **289717-75-5P**, Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-1-methyl- **289719-21-7P**, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-methyl-, hydrochloride
289719-22-8P, Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4-methoxy-, hydrochloride
 RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
 (preparation of phenoxybenzylamines as monoamine reuptake inhibitors)

RN 146520-69-6 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

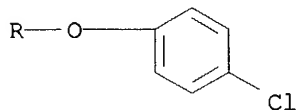
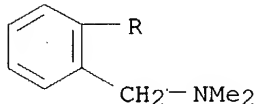
RN 146797-20-8 HCAPLUS

CN Benzenemethanamine, 2-(4-chlorophenoxy)-N,N-dimethyl-, (2Z)-2-butenedioate
(1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 146797-19-5

CMF C15 H16 Cl N O

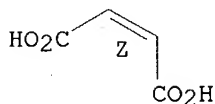


CM 2

CRN 110-16-7

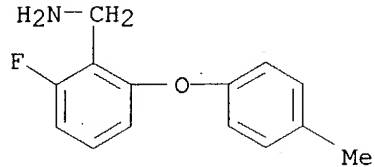
CMF C4 H4 O4

Double bond geometry as shown.



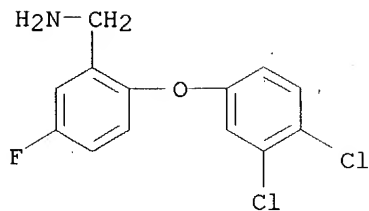
RN 289716-74-1 HCAPLUS

CN Benzenemethanamine, 2-fluoro-6-(4-methylphenoxy)- (9CI) (CA INDEX NAME)



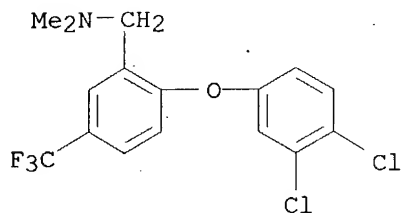
RN 289716-75-2 HCAPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro- (9CI) (CA INDEX NAME)



RN 289716-79-6 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-5-(trifluoromethyl)- (9CI) (CA INDEX NAME)



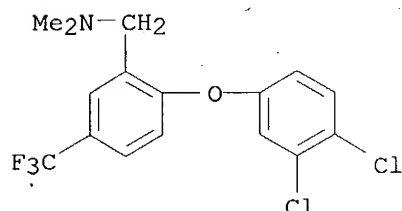
RN 289716-80-9 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-5-(trifluoromethyl)-, (2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 289716-79-6

CMF C16 H14 Cl2 F3 N O

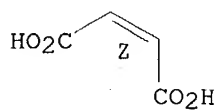


CM 2

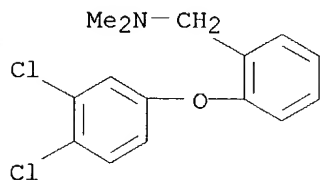
CRN 110-16-7

CMF C4 H4 O4

Double bond geometry as shown.



RN 289716-82-1 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-, hydrochloride
 (9CI) (CA INDEX NAME)

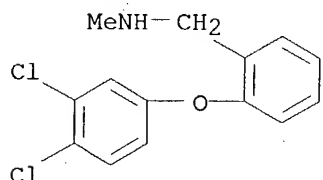


● HCl

RN 289716-85-4 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-, (2Z)-2-butenedioate
 (4:3) (9CI) (CA INDEX NAME)

CM 1

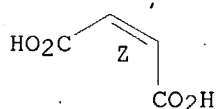
CRN 289716-84-3
 CMF C₁₄ H₁₃ Cl₂ N O



CM 2

CRN 110-16-7
 CMF C₄ H₄ O₄

Double bond geometry as shown.



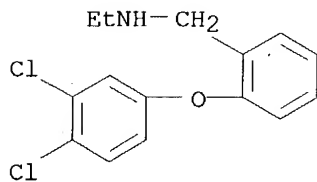
RN 289716-88-7 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-ethyl-, (2Z)-2-butenedioate
(1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 289716-87-6

CMF C15 H15 Cl2 N O

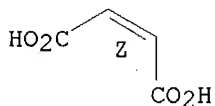


CM 2

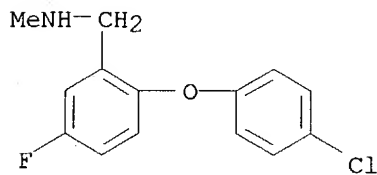
CRN 110-16-7

CMF C4 H4 O4

Double bond geometry as shown.



RN 289716-89-8 HCPLUS

CN Benzenemethanamine, 2-(4-chlorophenoxy)-5-fluoro-N-methyl-, hydrochloride
(9CI) (CA INDEX NAME)

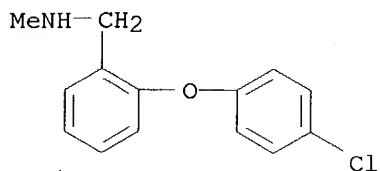
● HCl

RN 289716-91-2 HCPLUS

CN Benzenemethanamine, 2-(4-chlorophenoxy)-N-methyl-, (2Z)-2-butenedioate
(1:1) (9CI) (CA INDEX NAME)

CM 1

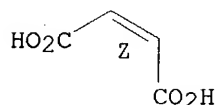
CRN 289716-90-1
 CMF C14 H14 Cl N O



CM 2

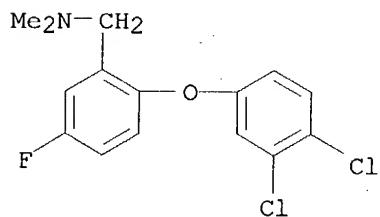
CRN 110-16-7
 CMF C4 H4 O4

Double bond geometry as shown.



RN 289716-92-3 HCPLUS

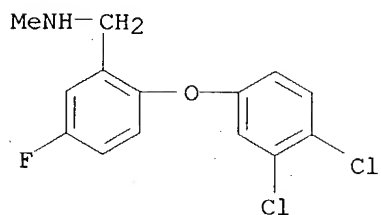
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N,N-dimethyl-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

RN 289716-93-4 HCPLUS

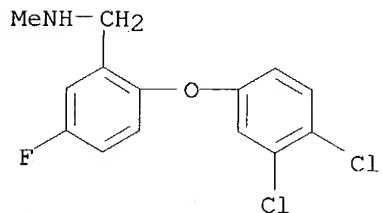
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N-methyl-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

RN 289716-94-5 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N-methyl- (9CI) (CA INDEX NAME)



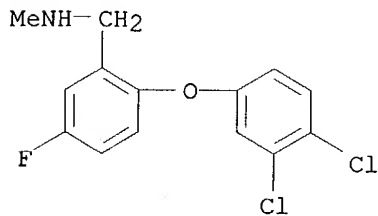
RN 289716-95-6 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N-methyl-, (2Z)-2-butenedioate (2:1) (9CI) (CA INDEX NAME)

CM 1

CRN 289716-94-5

CMF C14 H12 Cl2 F N O

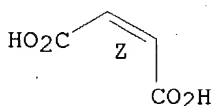


CM 2

CRN 110-16-7

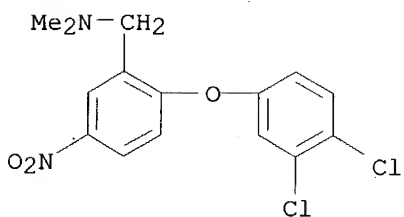
CMF C4 H4 O4

Double bond geometry as shown.



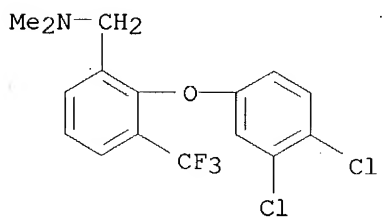
RN 289716-96-7 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-5-nitro- (9CI)
(CA INDEX NAME)



RN 289716-97-8 HCPLUS

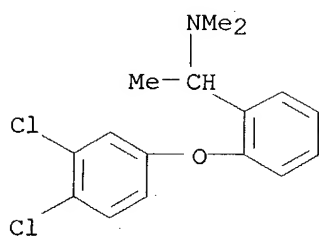
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-3-(trifluoromethyl)-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

RN 289716-98-9 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N, α -trimethyl-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

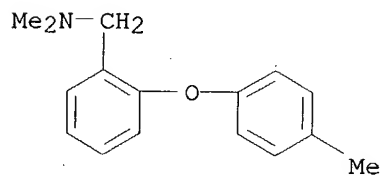
RN 289717-00-6 HCPLUS

CN Benzenemethanamine, N,N-dimethyl-2-(4-methylphenoxy)-, (2Z)-2-butenedioate
(1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 289716-99-0

CMF C16 H19 N O

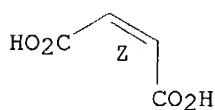


CM 2

CRN 110-16-7

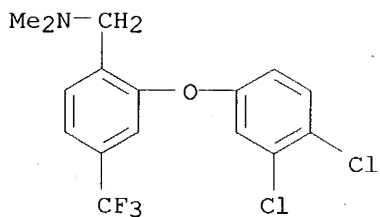
CMF C4 H4 O4

Double bond geometry as shown.



RN 289717-01-7 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-4-(trifluoromethyl)- (9CI) (CA INDEX NAME)



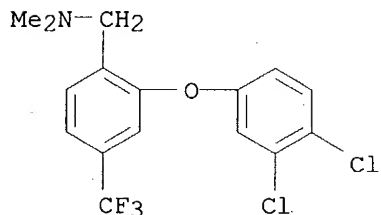
RN 289717-02-8 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-4-(trifluoromethyl)-, (2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 289717-01-7

CMF C16 H14 Cl2 F3 N O

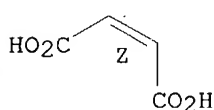


CM 2

CRN 110-16-7

CMF C4 H4 O4

Double bond geometry as shown.



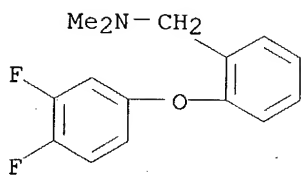
RN 289717-04-0 HCPLUS

CN Benzenemethanamine, 2-(3,4-difluorophenoxy)-N,N-dimethyl-, (2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 289717-03-9

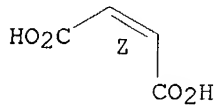
CMF C15 H15 F2 N O



CM 2

CRN 110-16-7
CMF C4 H4 O4

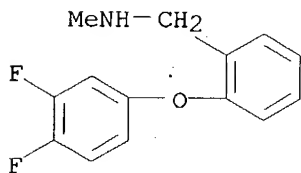
Double bond geometry as shown.



RN 289717-06-2 HCPLUS

CN Benzenemethanamine, 2-(3,4-difluorophenoxy)-N-methyl-, (2Z)-2-butenedioate
(1:1) (9CI) (CA INDEX NAME)

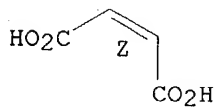
CM 1

CRN 289717-05-1
CMF C14 H13 F2 N O

CM 2

CRN 110-16-7
CMF C4 H4 O4

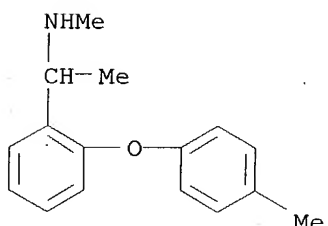
Double bond geometry as shown.



RN 289717-08-4 HCPLUS

CN Benzenemethanamine, N, α -dimethyl-2-(4-methylphenoxy)-, (2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

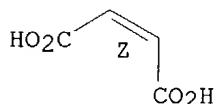
CM 1

CRN 289717-07-3
CMF C16 H19 N O

CM 2

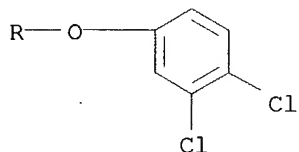
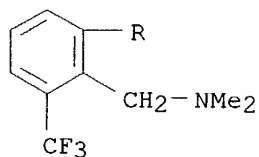
CRN 110-16-7
CMF C4 H4 O4

Double bond geometry as shown.



RN 289717-09-5 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-6-(trifluoromethyl)-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

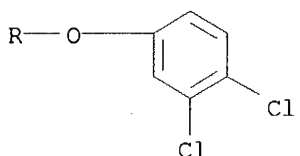
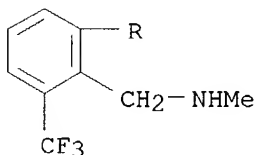
RN 289717-11-9 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-6-(trifluoromethyl)-, (2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 289717-10-8

CMF C15 H12 Cl2 F3 N O

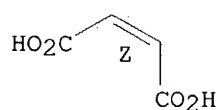


CM 2

CRN 110-16-7

CMF C4 H4 O4

Double bond geometry as shown.



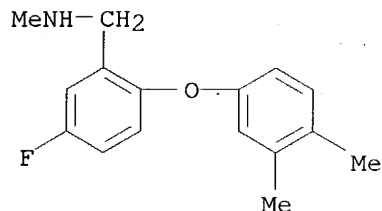
RN 289717-13-1 HCAPLUS

CN Benzenemethanamine, 2-(3,4-dimethylphenoxy)-5-fluoro-N-methyl-,
(2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 289717-12-0

CMF C16 H18 F N O

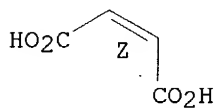


CM 2

CRN 110-16-7

CMF C4 H4 O4

Double bond geometry as shown.



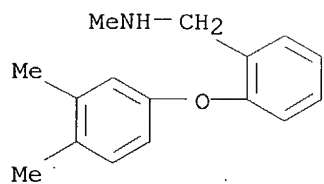
RN 289717-15-3 HCAPLUS

CN Benzenemethanamine, 2-(3,4-dimethylphenoxy)-N-methyl-, (2Z)-2-butenedioate
(1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 289717-14-2

CMF C16 H19 N O

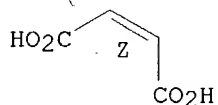


CM 2

CRN 110-16-7

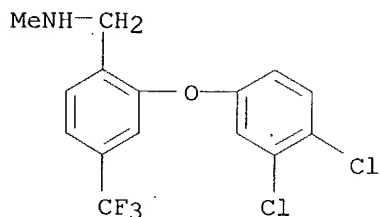
CMF C₈ H₁₂ O₄

Double bond geometry as shown.



RN 289717-16-4 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-4-(trifluoromethyl)-(9CI) (CA INDEX NAME)



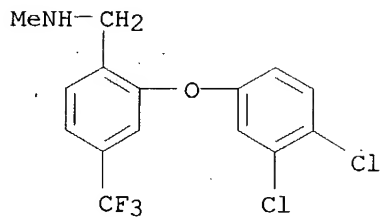
RN 289717-17-5 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-4-(trifluoromethyl)-(2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 289717-16-4

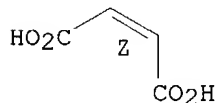
CMF C₁₅ H₁₂ Cl₂ F₃ N O



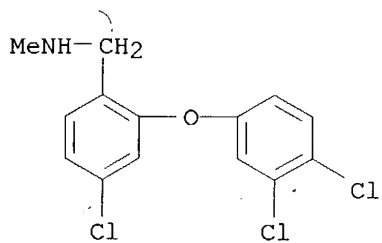
CM 2

CRN 110-16-7
CMF C₁₄ H₁₂ Cl₃ N O₄

Double bond geometry as shown.



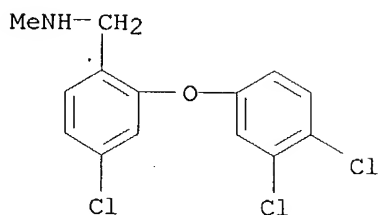
RN 289717-18-6 HCPLUS
 CN Benzenemethanamine, 4-chloro-2-(3,4-dichlorophenoxy)-N-methyl- (9CI) (CA INDEX NAME)



RN 289717-19-7 HCPLUS
 CN Benzenemethanamine, 4-chloro-2-(3,4-dichlorophenoxy)-N-methyl-, (2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

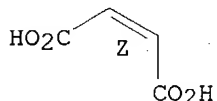
CRN 289717-18-6
CMF C₁₄ H₁₂ Cl₃ N O



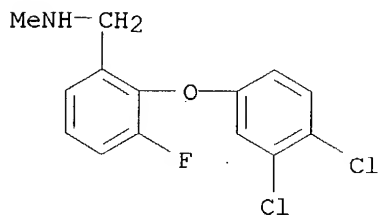
CM 2

CRN 110-16-7
CMF C₄ H₄ O₄

Double bond geometry as shown.

RN 289717-23-3 HCPLUS
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-3-fluoro-N-methyl-,
(2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

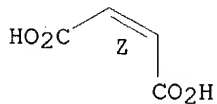
CM 1

CRN 289717-22-2
CMF C₁₄ H₁₂ Cl₂ F N O

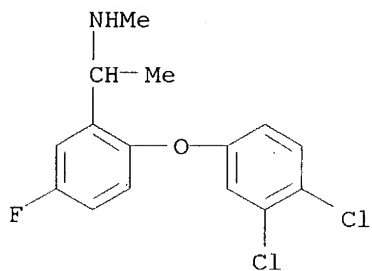
CM 2

CRN 110-16-7
CMF C₄ H₄ O₄

Double bond geometry as shown.



RN 289717-24-4 HCAPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N, α -dimethyl-
(9CI) (CA INDEX NAME)

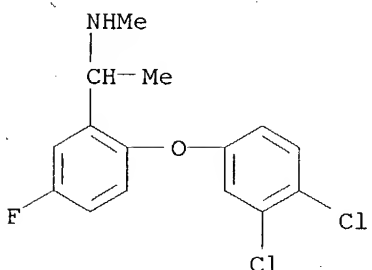
RN 289717-25-5 HCAPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N, α -dimethyl-,
(2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 289717-24-4

CMF C15 H14 Cl2 F N O

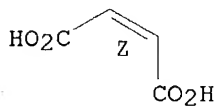


CM 2

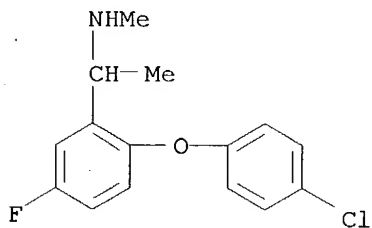
CRN 110-16-7

CMF C4 H4 O4

Double bond geometry as shown.



RN 289717-26-6 HCAPLUS

CN Benzenemethanamine, 2-(4-chlorophenoxy)-5-fluoro-N, α -dimethyl-, hydrochloride (9CI) (CA INDEX NAME)

● HCl

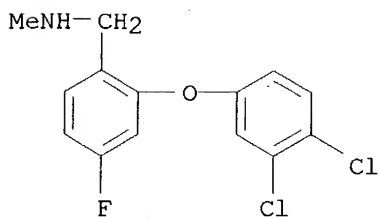
RN 289717-28-8 HCAPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4-fluoro-N-methyl-, (2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 289717-27-7

CMF C14 H12 Cl2 F N O

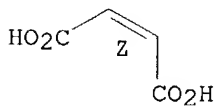


CM 2

CRN 110-16-7

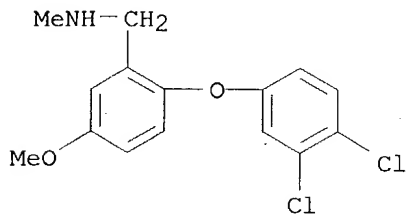
CMF C4 H4 O4

Double bond geometry as shown.



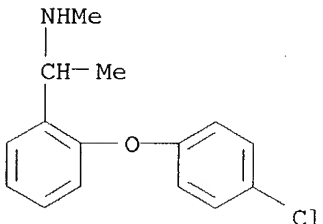
RN 289717-29-9 HCAPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-methoxy-N-methyl-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

RN 289717-30-2 HCAPLUS

CN Benzenemethanamine, 2-(4-chlorophenoxy)-N, α -dimethyl-, hydrochloride (9CI) (CA INDEX NAME)

● HCl

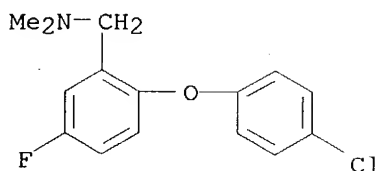
RN 289717-32-4 HCAPLUS

CN Benzenemethanamine, 2-(4-chlorophenoxy)-5-fluoro-N,N-dimethyl-, (2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 289717-31-3

CMF C15 H15 Cl F N O

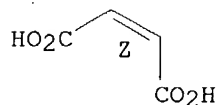


CM 2

CRN 110-16-7

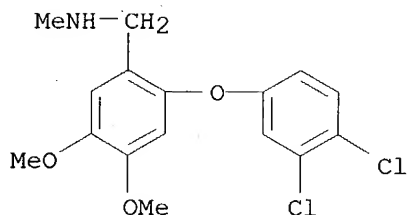
CMF C4 H4 O4

Double bond geometry as shown.



RN 289717-33-5 HCPLUS

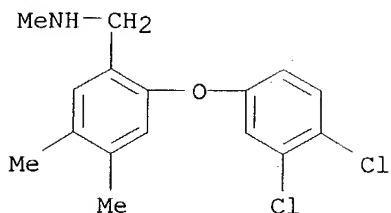
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4,5-dimethoxy-N-methyl-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

RN 289717-34-6 HCPLUS

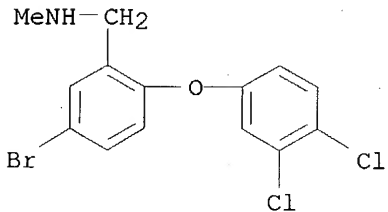
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,4,5-trimethyl-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

RN 289717-35-7 HCPLUS

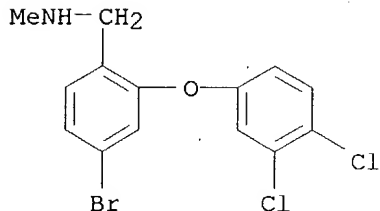
CN Benzenemethanamine, 5-bromo-2-(3,4-dichlorophenoxy)-N-methyl-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

RN 289717-36-8 HCPLUS

CN Benzenemethanamine, 4-bromo-2-(3,4-dichlorophenoxy)-N-methyl-, hydrochloride (9CI) (CA INDEX NAME)

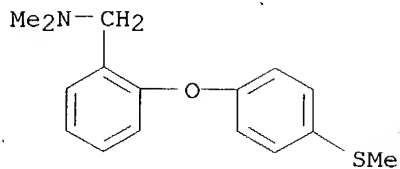


● HCl

RN 289717-37-9 HCPLUS

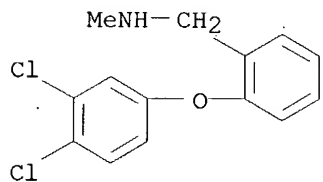
CN Benzenemethanamine, N,N-dimethyl-2-[4-(methylthio)phenoxy]-, hydrochloride

(9CI) (CA INDEX NAME)



● HCl

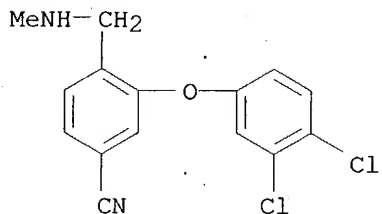
RN 289717-38-0 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-, hydrochloride (9CI)
(CA INDEX NAME)

● HCl

RN 289717-39-1 HCPLUS

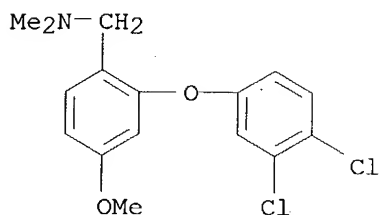
CN Benzonitrile, 3-(3,4-dichlorophenoxy)-4-[(methylamino)methyl]-, monohydrochloride (9CI) (CA INDEX NAME)



● HCl

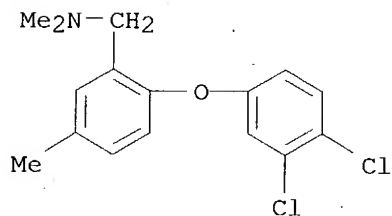
RN 289717-41-5 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4-methoxy-N,N-dimethyl-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

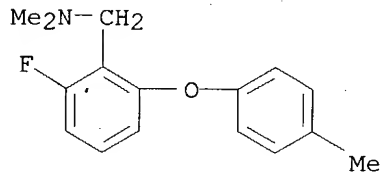
RN 289717-42-6 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N,5-trimethyl-,
hydrochloride (9CI) (CA INDEX NAME)

● HCl

RN 289717-43-7 HCPLUS

CN Benzenemethanamine, 2-fluoro-N,N-dimethyl-6-(4-methylphenoxy)- (9CI) (CA INDEX NAME)

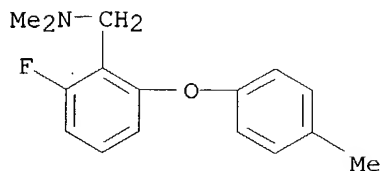


RN 289717-44-8 HCPLUS

CN Benzenemethanamine, 2-fluoro-N,N-dimethyl-6-(4-methylphenoxy)-,
(2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

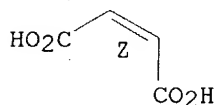
CRN 289717-43-7
 CMF C16 H18 F N O



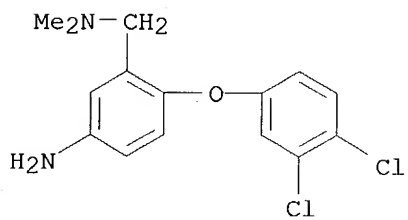
CM 2

CRN 110-16-7
 CMF C4 H4 O4

Double bond geometry as shown.

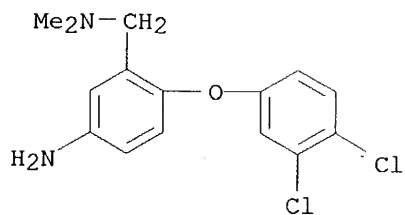


RN 289717-45-9 HCPLUS
 CN Benzenemethanamine, 5-amino-2-(3,4-dichlorophenoxy)-N,N-dimethyl-, monohydrochloride (9CI) (CA INDEX NAME)

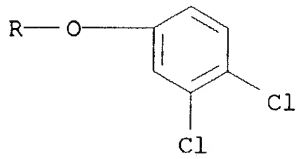
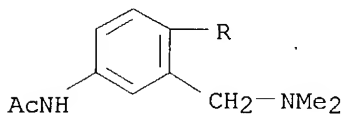


● HCl

RN 289717-46-0 HCPLUS
 CN Benzenemethanamine, 5-amino-2-(3,4-dichlorophenoxy)-N,N-dimethyl- (9CI) (CA INDEX NAME)

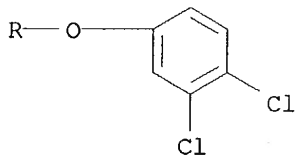
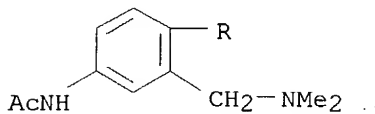


RN 289717-47-1 HCAPLUS
 CN Acetamide, N-[4-(3,4-dichlorophenoxy)-3-[(dimethylamino)methyl]phenyl]-, monohydrochloride (9CI) (CA INDEX NAME)



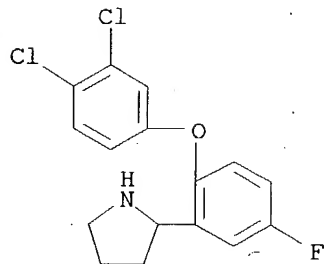
● HCl

RN 289717-48-2 HCAPLUS
 CN Acetamide, N-[4-(3,4-dichlorophenoxy)-3-[(dimethylamino)methyl]phenyl]- (9CI) (CA INDEX NAME)



RN 289717-49-3 HCAPLUS
 CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-, hydrochloride

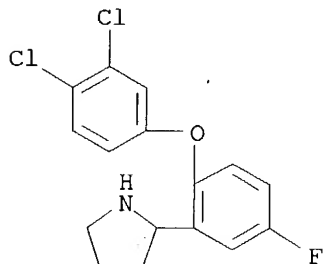
(9CI) (CA INDEX NAME)



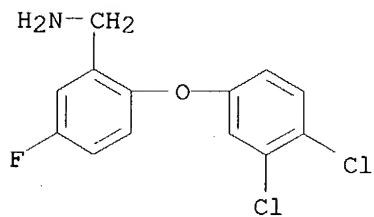
● HCl

RN 289717-50-6 HCPLUS

CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]- (9CI) (CA INDEX NAME)

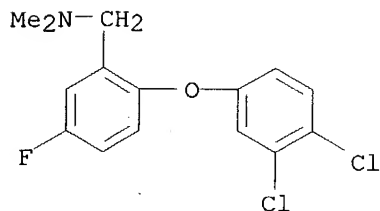


RN 289717-51-7 HCPLUS

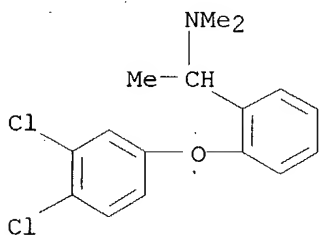
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-, hydrochloride (9CI)
(CA INDEX NAME)

● HCl

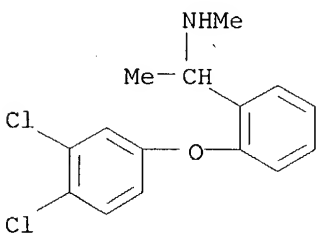
RN 289717-52-8 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N,N-dimethyl- (9CI)
(CA INDEX NAME)

RN 289717-53-9 HCPLUS

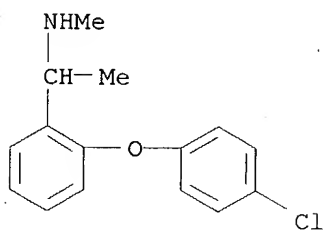
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N, α -trimethyl- (9CI)
(CA INDEX NAME)

RN 289717-54-0 HCPLUS

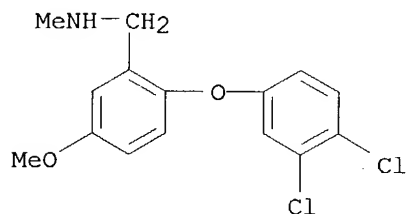
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N, α -dimethyl- (9CI) (CA
INDEX NAME)

RN 289717-55-1 HCPLUS

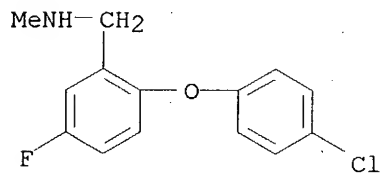
CN Benzenemethanamine, 2-(4-chlorophenoxy)-N, α -dimethyl- (9CI) (CA
INDEX NAME)



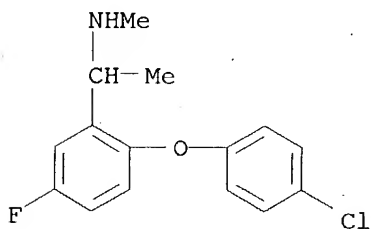
RN 289717-56-2 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-methoxy-N-methyl- (9CI) (CA INDEX NAME)



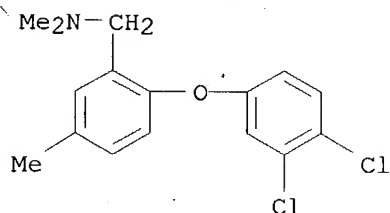
RN 289717-57-3 HCAPLUS
 CN Benzenemethanamine, 2-(4-chlorophenoxy)-5-fluoro-N-methyl- (9CI) (CA INDEX NAME)



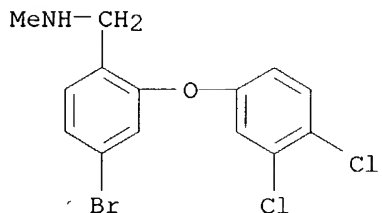
RN 289717-58-4 HCAPLUS
 CN Benzenemethanamine, 2-(4-chlorophenoxy)-5-fluoro-N, α -dimethyl- (9CI)
 (CA INDEX NAME)



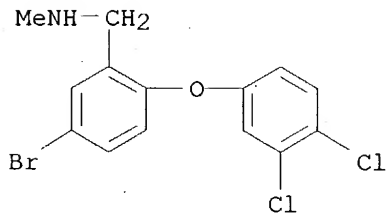
RN 289717-59-5 HCAPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N,5-trimethyl- (9CI) (CA
INDEX NAME)

RN 289717-60-8 HCAPLUS

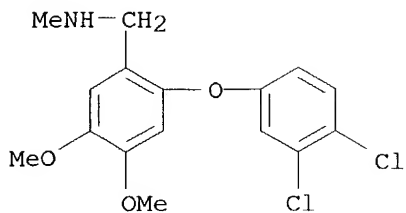
CN Benzenemethanamine, 4-bromo-2-(3,4-dichlorophenoxy)-N-methyl- (9CI) (CA
INDEX NAME)

RN 289717-61-9 HCAPLUS

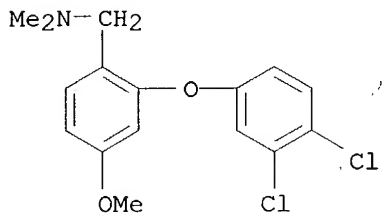
CN Benzenemethanamine, 5-bromo-2-(3,4-dichlorophenoxy)-N-methyl- (9CI) (CA
INDEX NAME)

RN 289717-62-0 HCAPLUS

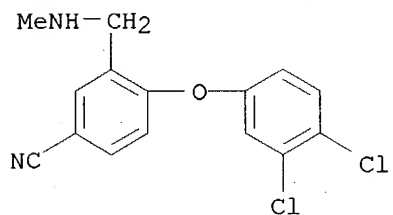
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4,5-dimethoxy-N-methyl- (9CI)
(CA INDEX NAME)



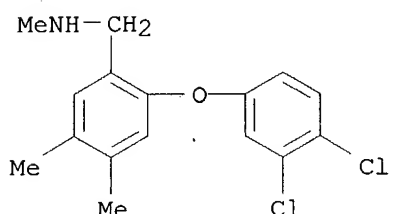
RN 289717-63-1 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4-methoxy-N,N-dimethyl- (9CI)
 (CA INDEX NAME)



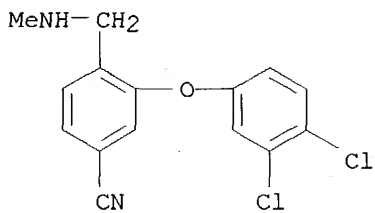
RN 289717-64-2 HCPLUS
 CN Benzonitrile, 4-(3,4-dichlorophenoxy)-3-[(methylamino)methyl]- (9CI) (CA INDEX NAME)



RN 289717-65-3 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,4,5-trimethyl- (9CI) (CA INDEX NAME)

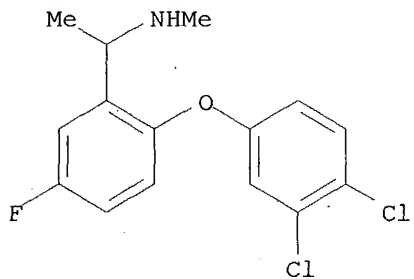


RN 289717-66-4 HCPLUS
 CN Benzonitrile, 3-(3,4-dichlorophenoxy)-4-[(methylamino)methyl]- (9CI) (CA INDEX NAME)



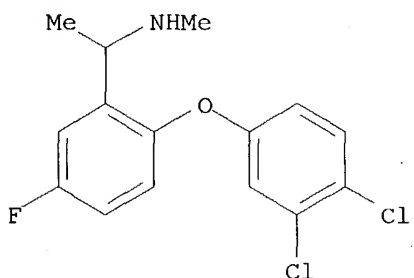
RN 289717-67-5 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N, α -dimethyl-, (+)- (9CI) (CA INDEX NAME)

Rotation (+).



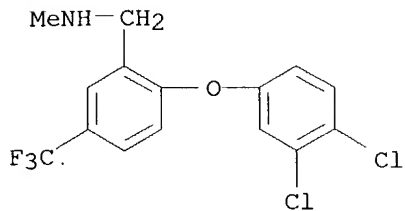
RN 289717-68-6 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N, α -dimethyl-, (-)- (9CI) (CA INDEX NAME)

Rotation (-).

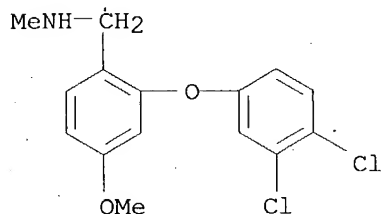


RN 289717-69-7 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-5-(trifluoromethyl)-

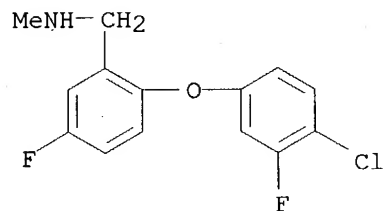
(9CI) (CA INDEX NAME)



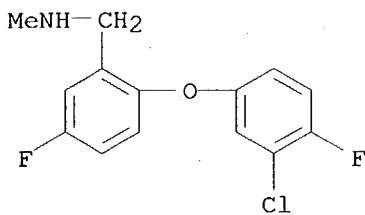
RN 289717-70-0 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4-methoxy-N-methyl- (9CI) (CA INDEX NAME)



RN 289717-71-1 HCPLUS
 CN Benzenemethanamine, 2-(4-chloro-3-fluorophenoxy)-5-fluoro-N-methyl- (9CI) (CA INDEX NAME)



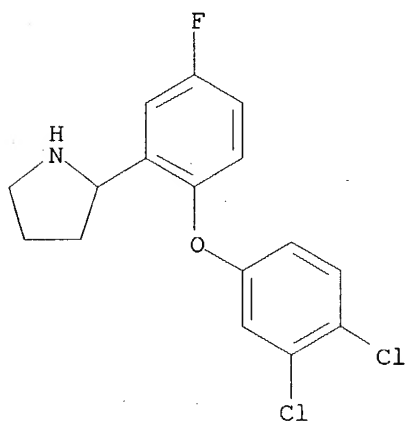
RN 289717-72-2 HCPLUS
 CN Benzenemethanamine, 2-(3-chloro-4-fluorophenoxy)-5-fluoro-N-methyl- (9CI) (CA INDEX NAME)



RN 289717-73-3 HCPLUS

CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-, (-)- (9CI) (CA INDEX NAME)

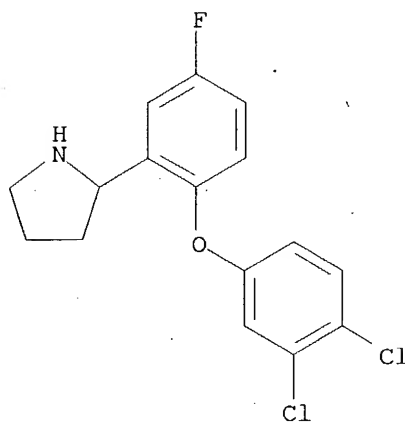
Rotation (-).



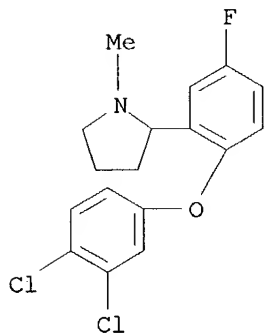
RN 289717-74-4 HCPLUS

CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-, (+)- (9CI) (CA INDEX NAME)

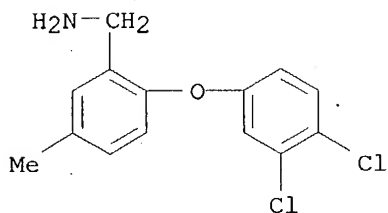
Rotation (+).



RN 289717-75-5 HCAPLUS
 CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-1-methyl- (9CI)
 (CA INDEX NAME)

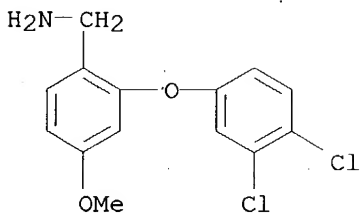


RN 289719-21-7 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-methyl-, hydrochloride (9CI)
 (CA INDEX NAME)



● HCl

RN 289719-22-8 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4-methoxy-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

L51 ANSWER 12 OF 44 HCPLUS COPYRIGHT 2004 ACS on STN
 ACCESSION NUMBER: 2002:674788 HCPLUS
 DOCUMENT NUMBER: 137:195595
 TITLE: Atypical antipsychotic-antidepressant combination for treatment of depression, obsessive compulsive disorder, and psychosis
 INVENTOR(S): Howard, Harry R., Jr.
 PATENT ASSIGNEE(S): Pfizer Inc., USA
 SOURCE: U.S. Pat. Appl. Publ., 20 pp.
 CODEN: USXXCO
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2002123490	A1	20020905	US 2001-10651	20011206
EP 1238676	A1	20020911	EP 2002-251153	20020220
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR				
JP 2002308801	A2	20021023	JP 2002-50579	20020227
PRIORITY APPLN. INFO.:			US 2001-272619P	P 20010301

OTHER SOURCE(S): MARPAT 137:195595

AB The invention provides a method for treating depression, obsessive compulsive disorder, and psychosis in a mammal, including a human, by administering to the mammal an atypical antipsychotic in combination with an antidepressant agent with improvement in efficiency. It also provides pharmaceutical compns. containing a pharmaceutically acceptable carrier, an atypical antipsychotic; and a serotonin reuptake inhibitor.

IT 289716-79-6 289716-94-5 289717-01-7
 289717-16-4 289717-18-6 289717-24-4
 289717-48-2 289717-50-6 289717-52-8
 289717-53-9 289717-54-0 289717-55-1
 289717-56-2 289717-57-3 289717-58-4
 289717-59-5 289717-60-8 289717-61-9
 289717-62-0 289717-63-1 289717-64-2
 289717-65-3 289717-66-4 289717-67-5
 289717-68-6 289717-69-7 289717-70-0
 289717-71-1 289717-72-2 289717-73-3
 289717-74-4 289717-75-5 444888-21-5

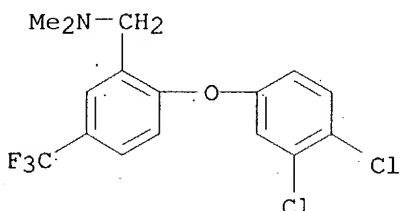
444888-23-7 444888-24-8 444888-25-9
 444888-27-1 444888-28-2 444888-29-3
 444888-30-6 444888-31-7 444888-32-8
 444888-33-9 444888-34-0 444888-35-1
 444888-36-2 444888-37-3 444888-38-4
 444888-39-5 444888-40-8 444888-41-9
 444888-42-0 444888-43-1 444888-45-3
 444888-46-4 444888-49-7 454456-38-3
 454456-43-0 454456-66-7 454456-75-8
 454473-98-4

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL
 (Biological study); USES (Uses)

(atypical antipsychotic-antidepressant combination for treatment of
 depression, obsessive compulsive disorder, and psychosis)

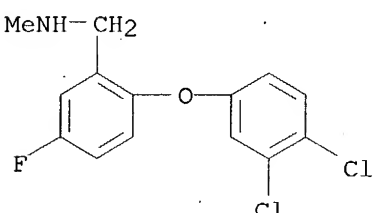
RN 289716-79-6 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-5-
 (trifluoromethyl)- (9CI) (CA INDEX NAME)



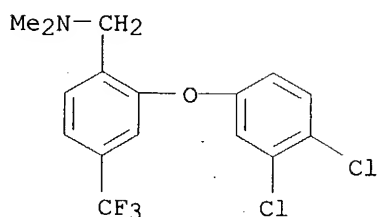
RN 289716-94-5 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N-methyl- (9CI) (CA
 INDEX NAME)



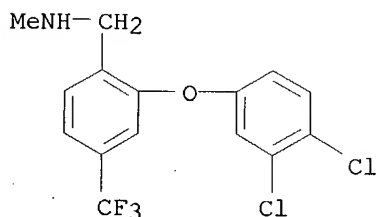
RN 289717-01-7 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-4-
 (trifluoromethyl)- (9CI) (CA INDEX NAME)



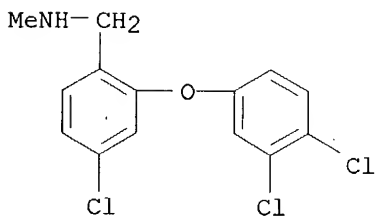
RN 289717-16-4 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-4-(trifluoromethyl)-(9CI) (CA INDEX NAME)



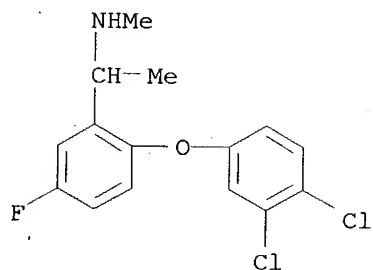
RN 289717-18-6 HCPLUS

CN Benzenemethanamine, 4-chloro-2-(3,4-dichlorophenoxy)-N-methyl- (9CI) (CA INDEX NAME)



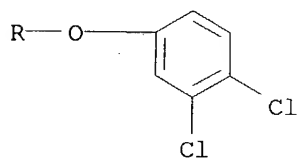
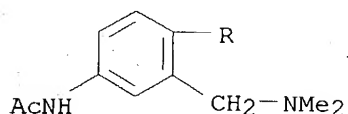
RN 289717-24-4 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N, α -dimethyl-(9CI) (CA INDEX NAME)



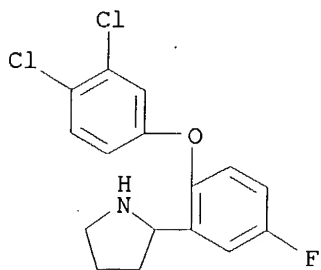
RN 289717-48-2 HCPLUS

CN Acetamide, N-[4-(3,4-dichlorophenoxy)-3-[(dimethylamino)methyl]phenyl]- (9CI) (CA INDEX NAME)



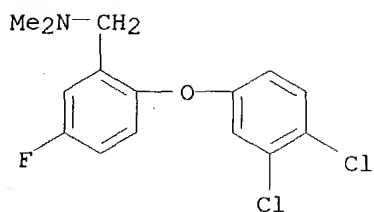
RN 289717-50-6 HCPLUS

CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]- (9CI) (CA INDEX NAME)

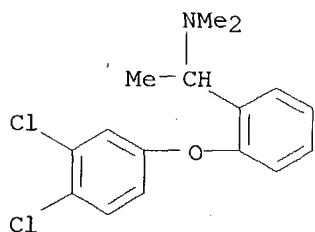


RN 289717-52-8 HCPLUS

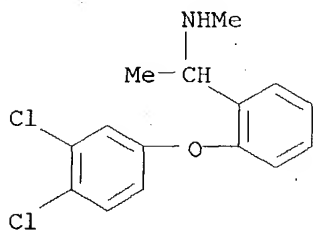
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N,N-dimethyl- (9CI) (CA INDEX NAME)



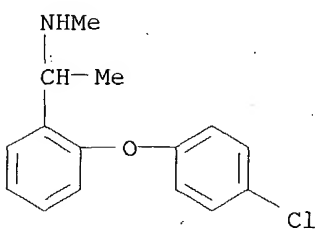
RN 289717-53-9 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N, α -trimethyl- (9CI)
 (CA INDEX NAME)



RN 289717-54-0 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N, α -dimethyl- (9CI) (CA
 INDEX NAME)

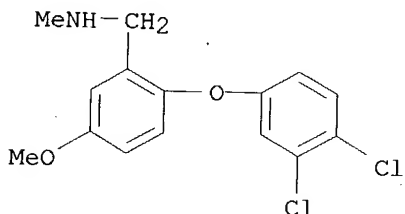


RN 289717-55-1 HCPLUS
 CN Benzenemethanamine, 2-(4-chlorophenoxy)-N, α -dimethyl- (9CI) (CA
 INDEX NAME)



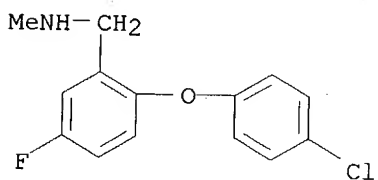
RN 289717-56-2 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-methoxy-N-methyl- (9CI) (CA INDEX NAME)

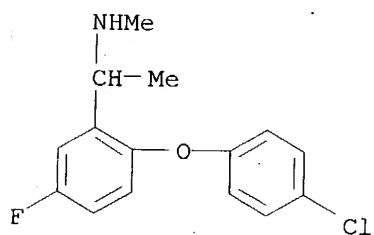


RN 289717-57-3 HCPLUS

CN Benzenemethanamine, 2-(4-chlorophenoxy)-5-fluoro-N-methyl- (9CI) (CA INDEX NAME)

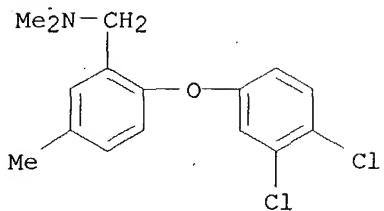


RN 289717-58-4 HCPLUS

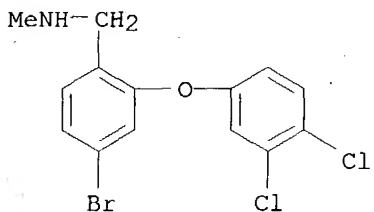
CN Benzenemethanamine, 2-(4-chlorophenoxy)-5-fluoro-N, α -dimethyl- (9CI)
(CA INDEX NAME)

RN 289717-59-5 HCPLUS

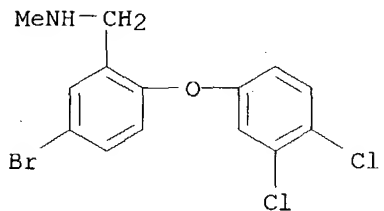
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N,5-trimethyl- (9CI) (CA INDEX NAME)



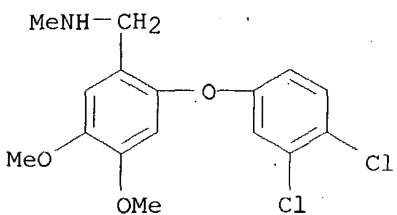
RN 289717-60-8 HCAPLUS
 CN Benzenemethanamine, 4-bromo-2-(3,4-dichlorophenoxy)-N-methyl- (9CI) (CA INDEX NAME)



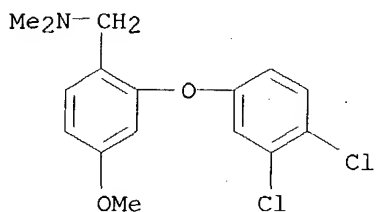
RN 289717-61-9 HCAPLUS
 CN Benzenemethanamine, 5-bromo-2-(3,4-dichlorophenoxy)-N-methyl- (9CI) (CA INDEX NAME)



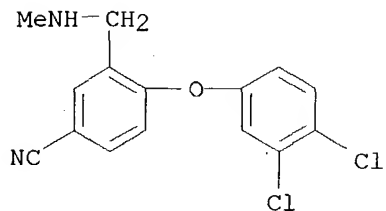
RN 289717-62-0 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4,5-dimethoxy-N-methyl- (9CI) (CA INDEX NAME)



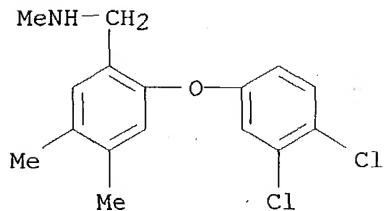
RN 289717-63-1 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4-methoxy-N,N-dimethyl- (9CI)
 (CA INDEX NAME)



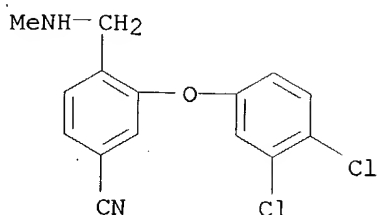
RN 289717-64-2 HCPLUS
 CN Benzonitrile, 4-(3,4-dichlorophenoxy)-3-[(methylamino)methyl]- (9CI) (CA INDEX NAME)



RN 289717-65-3 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,4,5-trimethyl- (9CI) (CA INDEX NAME)



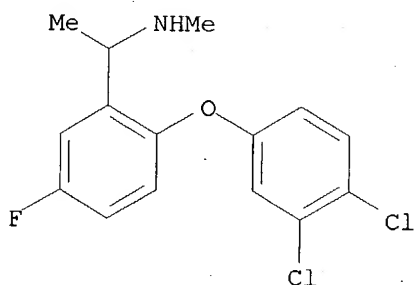
RN 289717-66-4 HCPLUS
 CN Benzonitrile, 3-(3,4-dichlorophenoxy)-4-[(methylamino)methyl]- (9CI) (CA INDEX NAME)



RN 289717-67-5 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N, α -dimethyl-,
(+)- (9CI) (CA INDEX NAME)

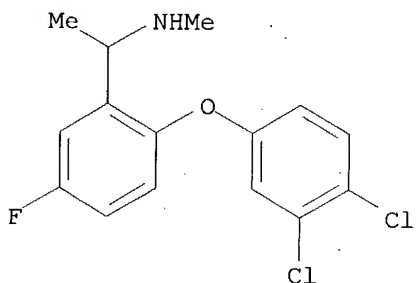
Rotation (+).



RN 289717-68-6 HCPLUS

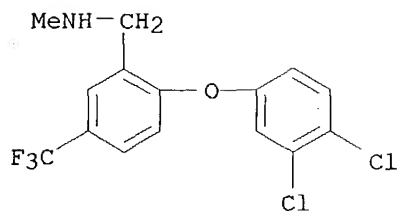
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N, α -dimethyl-,
(-)- (9CI) (CA INDEX NAME)

Rotation (-).

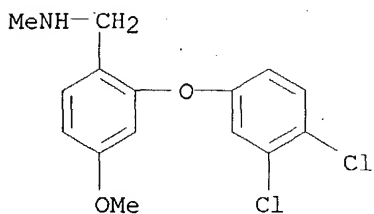


RN 289717-69-7 HCPLUS

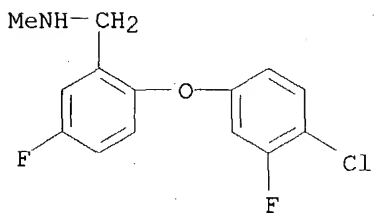
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-5-(trifluoromethyl)-
(9CI) (CA INDEX NAME)



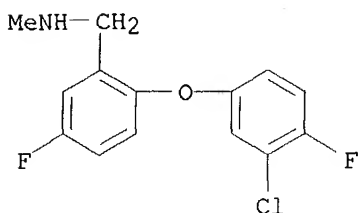
RN 289717-70-0 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4-methoxy-N-methyl- (9CI) (CA INDEX NAME)



RN 289717-71-1 HCPLUS
 CN Benzenemethanamine, 2-(4-chloro-3-fluorophenoxy)-5-fluoro-N-methyl- (9CI) (CA INDEX NAME)



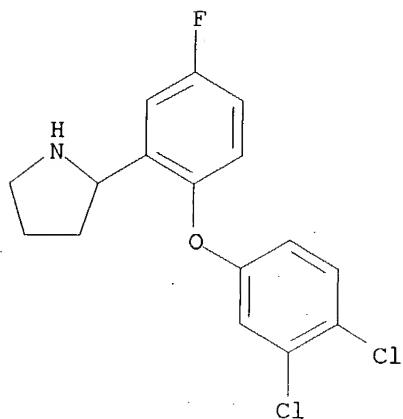
RN 289717-72-2 HCPLUS
 CN Benzenemethanamine, 2-(3-chloro-4-fluorophenoxy)-5-fluoro-N-methyl- (9CI) (CA INDEX NAME)



RN 289717-73-3 HCPLUS

CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-, (-)- (9CI) (CA
INDEX NAME)

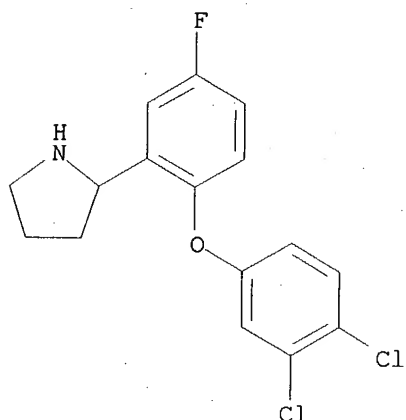
Rotation (-).



RN 289717-74-4 HCPLUS

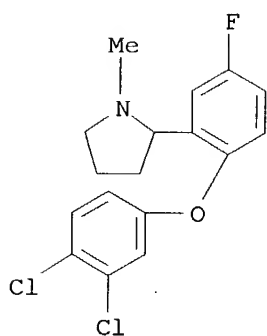
CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-, (+)- (9CI) (CA
INDEX NAME)

Rotation (+).



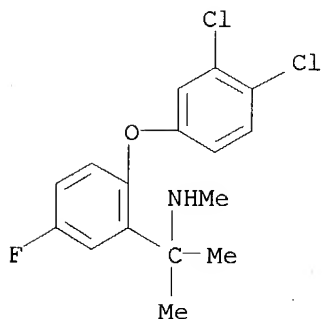
RN 289717-75-5 HCPLUS

CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-1-methyl- (9CI)
(CA INDEX NAME)



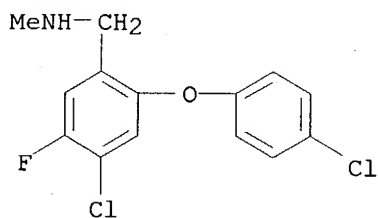
RN 444888-21-5 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N,α,α-trimethyl- (9CI) (CA INDEX NAME)



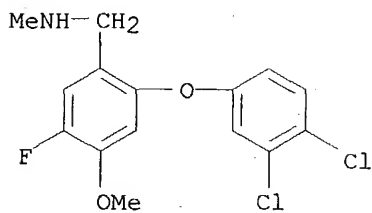
RN 444888-23-7 HCPLUS

CN Benzenemethanamine, 4-chloro-2-(4-chlorophenoxy)-5-fluoro-N-methyl- (9CI) (CA INDEX NAME)

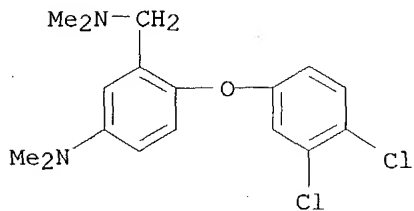


RN 444888-24-8 HCPLUS

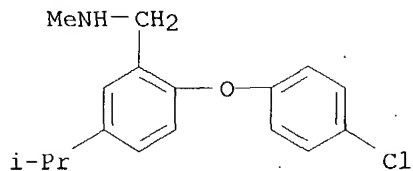
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-4-methoxy-N-methyl- (9CI) (CA INDEX NAME)



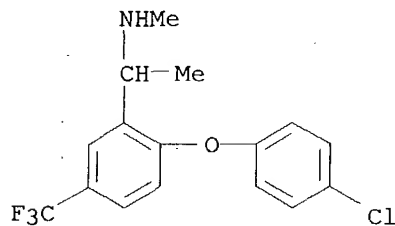
RN 444888-25-9 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-(dimethylamino)-N,N-dimethyl-
 (9CI) (CA INDEX NAME)



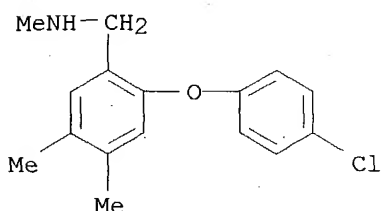
RN 444888-27-1 HCAPLUS
 CN Benzenemethanamine, 2-(4-chlorophenoxy)-N-methyl-5-(1-methylethyl)- (9CI)
 (CA INDEX NAME)



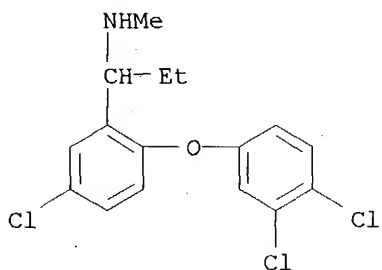
RN 444888-28-2 HCAPLUS
 CN Benzenemethanamine, 2-(4-chlorophenoxy)-N, α -dimethyl-5-
 (trifluoromethyl)- (9CI) (CA INDEX NAME)



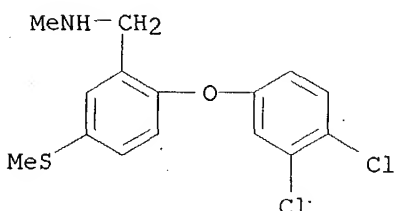
RN 444888-29-3 HCPLUS
 CN Benzenemethanamine, 2-(4-chlorophenoxy)-N,4,5-trimethyl- (9CI) (CA INDEX NAME)



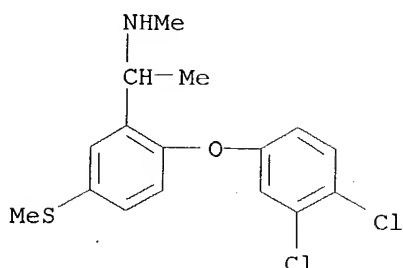
RN 444888-30-6 HCPLUS
 CN Benzenemethanamine, 5-chloro-2-(3,4-dichlorophenoxy)- α -ethyl-N-methyl- (9CI) (CA INDEX NAME)



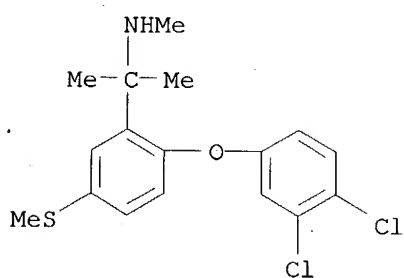
RN 444888-31-7 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-5-(methylthio)- (9CI)
 (CA INDEX NAME)



RN 444888-32-8 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N, α -dimethyl-5-(methylthio)- (9CI) (CA INDEX NAME)

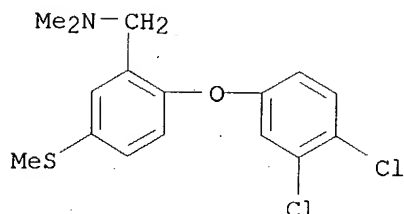


RN 444888-33-9 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N, α , α -trimethyl-5-(methylthio)- (9CI) (CA INDEX NAME)

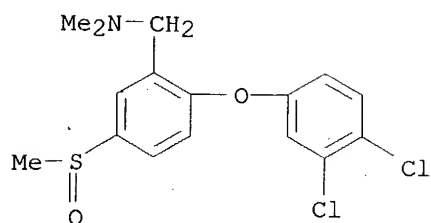
RN 444888-34-0 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-5-(methylthio)- (9CI) (CA INDEX NAME)

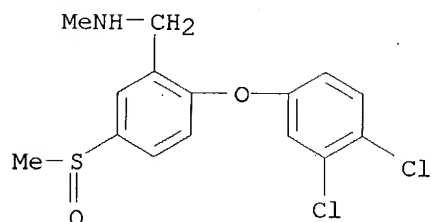


RN 444888-35-1 HCPLUS

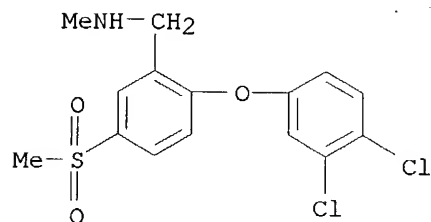
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-5-(methylsulfinyl)- (9CI) (CA INDEX NAME)



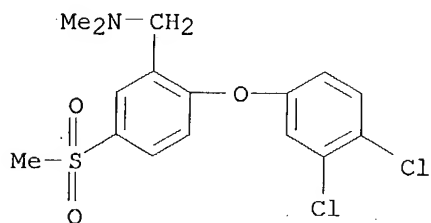
RN 444888-36-2 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-5-(methylsulfinyl)-
 (9CI) (CA INDEX NAME)



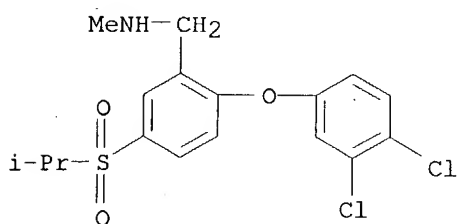
RN 444888-37-3 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-5-(methylsulfonyl)-
 (9CI) (CA INDEX NAME)



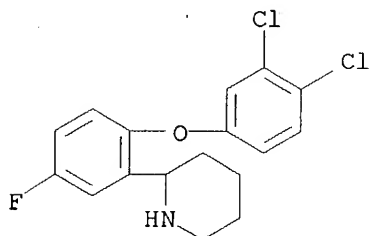
RN 444888-38-4 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-5-
 (methylsulfonyl)- (9CI) (CA INDEX NAME)



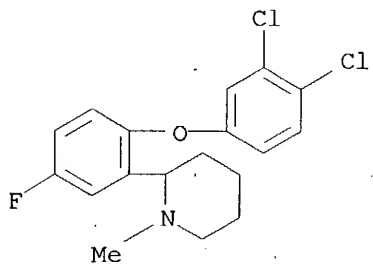
RN 444888-39-5 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-5-[(1-methylethyl)sulfonyl]- (9CI) (CA INDEX NAME)



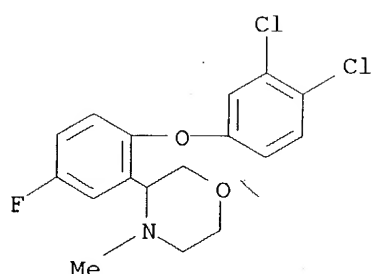
RN 444888-40-8 HCPLUS
 CN Piperidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]- (9CI) (CA INDEX NAME)



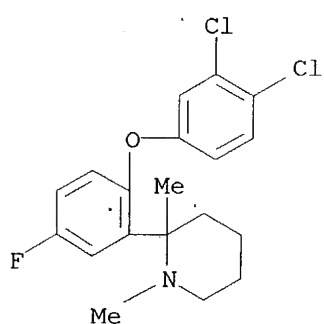
RN 444888-41-9 HCPLUS
 CN Piperidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-1-methyl- (9CI)
 (CA INDEX NAME)



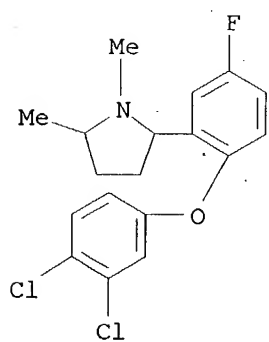
RN 444888-42-0 HCPLUS
 CN Morpholine, 3-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-4-methyl- (9CI)
 (CA INDEX NAME)



RN 444888-43-1 HCPLUS

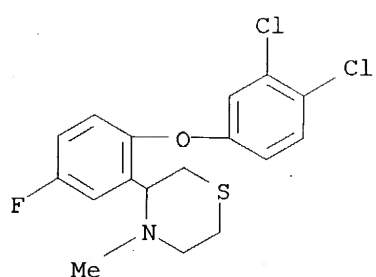
CN Piperidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-1,2-dimethyl- (9CI)
(CA INDEX NAME)

RN 444888-45-3 HCPLUS

CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-1,5-dimethyl-
(9CI) (CA INDEX NAME)

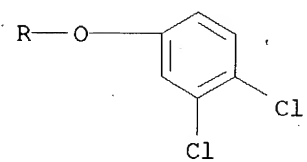
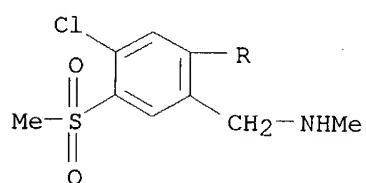
RN 444888-46-4 HCPLUS

CN Thiomorpholine, 3-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-4-methyl- (9CI)
(CA INDEX NAME)

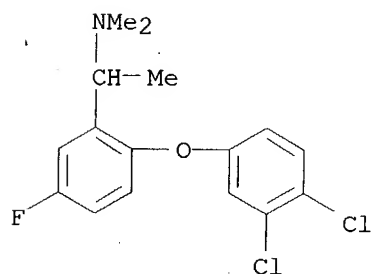


RN 444888-49-7 HCAPLUS

CN Benzenemethanamine, 4-chloro-2-(3,4-dichlorophenoxy)-N-methyl-5-(methylsulfonyl)- (9CI) (CA INDEX NAME)

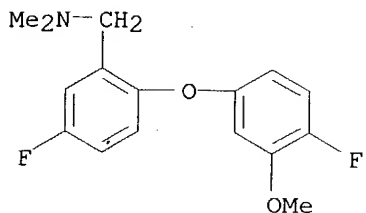


RN 454456-38-3 HCAPLUS

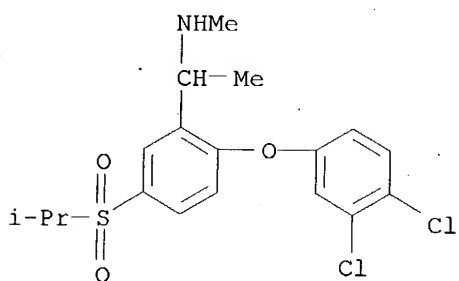
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N,N, α -trimethyl- (9CI) (CA INDEX NAME)

RN 454456-43-0 HCAPLUS

CN Benzenemethanamine, 5-fluoro-2-(4-fluoro-3-methoxyphenoxy)-N,N-dimethyl- (9CI) (CA INDEX NAME)

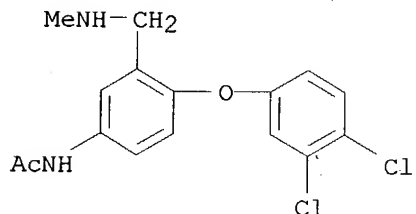


RN 454456-66-7 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N, α -dimethyl-5-[(1-methylethyl)sulfonyl]- (9CI) (CA INDEX NAME)

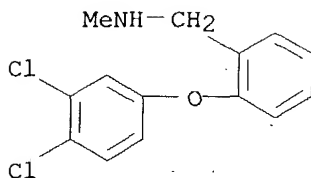
RN 454456-75-8 HCPLUS

CN Acetamide, N-[4-(3,4-dichlorophenoxy)-3-[(methylamino)methyl]phenyl]- (9CI) (CA INDEX NAME)



RN 454473-98-4 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-ar-fluoro-N-methyl- (9CI) (CA INDEX NAME)



D1-F

L51 ANSWER 13 OF 44 HCPLUS COPYRIGHT 2004 ACS on STN
 ACCESSION NUMBER: 2002:595509 HCPLUS
 DOCUMENT NUMBER: 137:135106
 TITLE: Combination of a 5-HT3 receptor antagonist with a serotonin reuptake inhibitor for the treatment of depression
 INVENTOR(S): Howard, Harry R.
 PATENT ASSIGNEE(S): USA
 SOURCE: U.S. Pat. Appl. Publ., 20 pp.
 CODEN: USXXCO
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2002107244	A1	20020808	US 2001-2303	20011102
EP 1230921	A1	20020814	EP 2002-250541	20020128
	R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR			
JP 2002275097	A2	20020925	JP 2002-20186	20020129
BR 2002000246	A	20021029	BR 2002-246	20020131

PRIORITY APPLN. INFO.: US 2001-266340P P 20010202

OTHER SOURCE(S): MARPAT 137:135106

AB The present invention relates to a method of treating depression or anxiety in a mammal, including a human, by administering to the mammal a 5-HT3 receptor antagonist in combination with a serotonin reuptake inhibitor (SRI) antidepressant agent with improvement in sexual function and/or reduction in gastro-intestinal side effects. It also relates to pharmaceutical compns. containing a pharmaceutically acceptable carrier, a 5-HT3 receptor antagonist and an SRI antidepressant. The ratio of the 5-HT3 receptor antagonist and the SRI antidepressant agent is between 0.001 to 1 and 1000 to 1, and especially between 0.01 to 1 and 100 to 1 (no data).

IT 289716-79-6 289716-94-5 289717-01-7
 289717-16-4 289717-18-6 289717-24-4
 289717-48-2 289717-50-6 289717-52-8
 289717-53-9 289717-54-0 289717-55-1
 289717-56-2 289717-57-3 289717-58-4
 289717-59-5 289717-60-8 289717-61-9
 289717-62-0 289717-63-1 289717-64-2
 289717-65-3 289717-66-4 289717-67-5

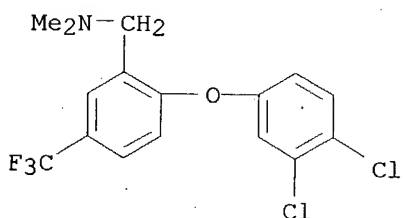
289717-68-6 289717-69-7 289717-70-0
 289717-71-1 289717-72-2 289717-73-3
 289717-74-4 289717-75-5 444888-21-5
 444888-22-6 444888-23-7 444888-24-8
 444888-25-9 444888-26-0 444888-27-1
 444888-28-2 444888-29-3 444888-30-6
 444888-31-7 444888-32-8 444888-33-9
 444888-34-0 444888-35-1 444888-36-2
 444888-37-3 444888-38-4 444888-39-5
 444888-40-8 444888-41-9 444888-42-0
 444888-43-1 444888-45-3 444888-46-4
 444888-48-6 444888-49-7

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(combination of 5-HT3 receptor antagonist with serotonin reuptake inhibitor for treatment of depression)

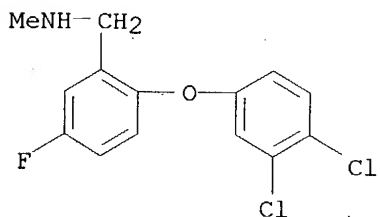
RN 289716-79-6 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-5-(trifluoromethyl)- (9CI) (CA INDEX NAME)



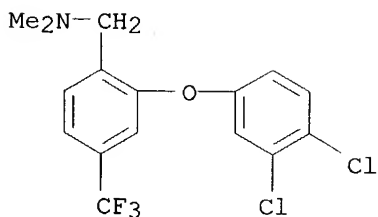
RN 289716-94-5 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N-methyl- (9CI) (CA INDEX NAME)

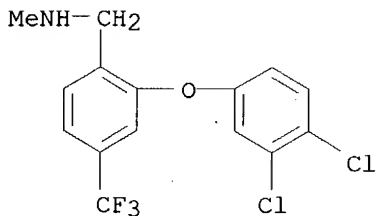


RN 289717-01-7 HCPLUS

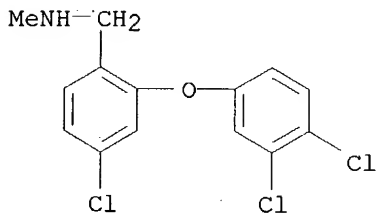
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-4-(trifluoromethyl)- (9CI) (CA INDEX NAME)



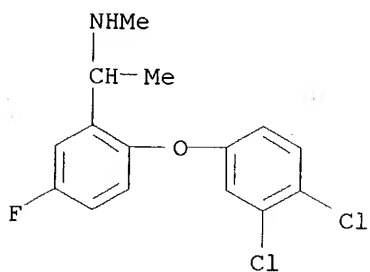
RN 289717-16-4 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-4-(trifluoromethyl)-
 (9CI) (CA INDEX NAME)



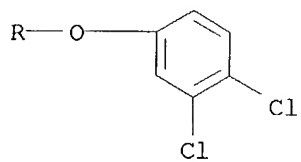
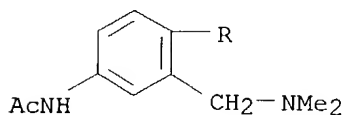
RN 289717-18-6 HCAPLUS
 CN Benzenemethanamine, 4-chloro-2-(3,4-dichlorophenoxy)-N-methyl- (9CI) (CA INDEX NAME)



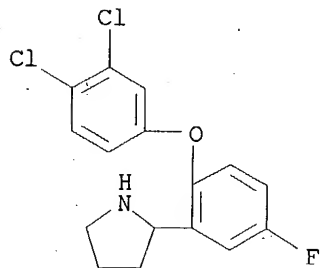
RN 289717-24-4 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N,α-dimethyl-
 (9CI) (CA INDEX NAME)



RN 289717-48-2 HCPLUS

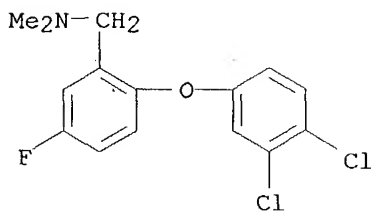
CN Acetamide, N-[4-(3,4-dichlorophenoxy)-3-(dimethylamino)methyl]phenyl-
(9CI) (CA INDEX NAME)

RN 289717-50-6 HCPLUS

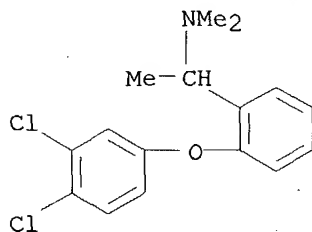
CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]- (9CI) (CA INDEX
NAME)

RN 289717-52-8 HCPLUS

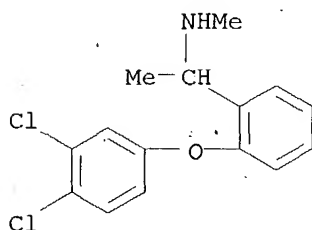
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N,N-dimethyl- (9CI)
(CA INDEX NAME)



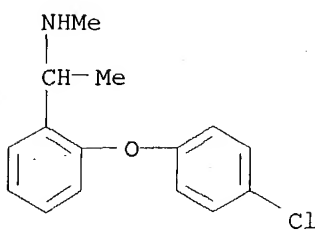
RN 289717-53-9 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N, α -trimethyl- (9CI)
 (CA INDEX NAME)



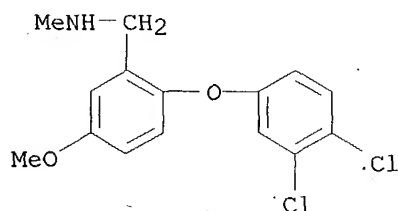
RN 289717-54-0 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N, α -dimethyl- (9CI) (CA
 INDEX NAME)



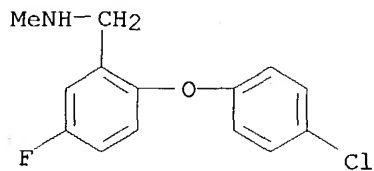
RN 289717-55-1 HCAPLUS
 CN Benzenemethanamine, 2-(4-chlorophenoxy)-N, α -dimethyl- (9CI) (CA
 INDEX NAME)



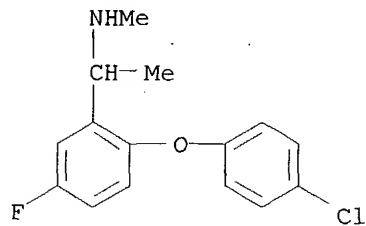
RN 289717-56-2 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-methoxy-N-methyl- (9CI) (CA INDEX NAME)



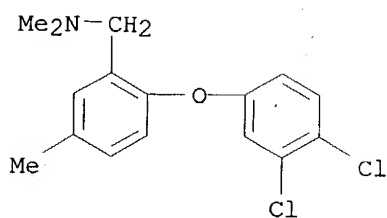
RN 289717-57-3 HCPLUS
 CN Benzenemethanamine, 2-(4-chlorophenoxy)-5-fluoro-N-methyl- (9CI) (CA INDEX NAME)



RN 289717-58-4 HCPLUS
 CN Benzenemethanamine, 2-(4-chlorophenoxy)-5-fluoro-N, α -dimethyl- (9CI) (CA INDEX NAME)

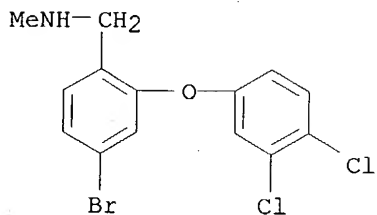


RN 289717-59-5 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N,5-trimethyl- (9CI) (CA INDEX NAME)



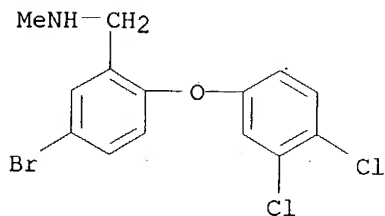
RN 289717-60-8 HCAPLUS

CN Benzenemethanamine, 4-bromo-2-(3,4-dichlorophenoxy)-N-methyl- (9CI) (CA INDEX NAME)



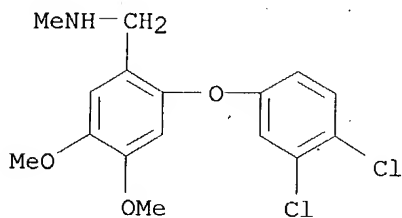
RN 289717-61-9 HCAPLUS

CN Benzenemethanamine, 5-bromo-2-(3,4-dichlorophenoxy)-N-methyl- (9CI) (CA INDEX NAME)

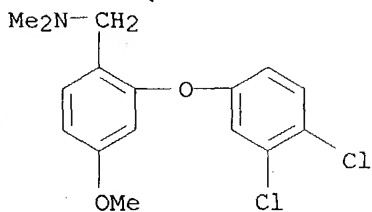


RN 289717-62-0 HCAPLUS

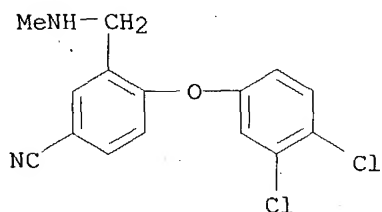
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4,5-dimethoxy-N-methyl- (9CI) (CA INDEX NAME)



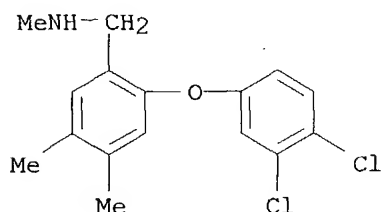
RN 289717-63-1 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4-methoxy-N,N-dimethyl- (9CI)
 (CA INDEX NAME)



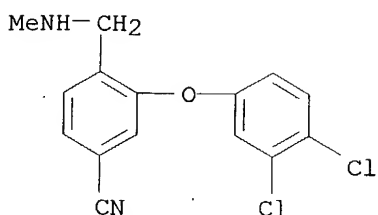
RN 289717-64-2 HCPLUS
 CN Benzonitrile, 4-(3,4-dichlorophenoxy)-3-[{methylamino)methyl]- (9CI) (CA INDEX NAME)



RN 289717-65-3 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,4,5-trimethyl- (9CI) (CA INDEX NAME)



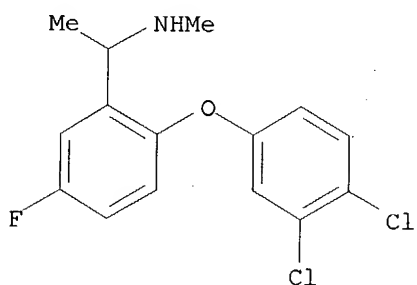
RN 289717-66-4 HCPLUS
 CN Benzonitrile, 3-(3,4-dichlorophenoxy)-4-[{methylamino)methyl]- (9CI) (CA INDEX NAME)



RN 289717-67-5 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N, α -dimethyl-,
(+)- (9CI) (CA INDEX NAME)

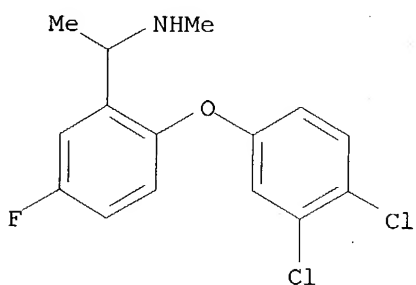
Rotation (+).



RN 289717-68-6 HCPLUS

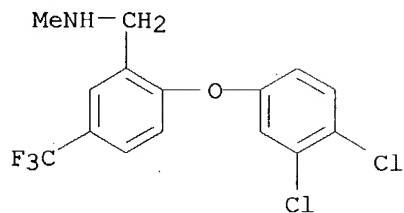
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N, α -dimethyl-,
(-)- (9CI) (CA INDEX NAME)

Rotation (-).

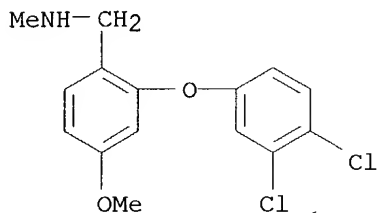


RN 289717-69-7 HCPLUS

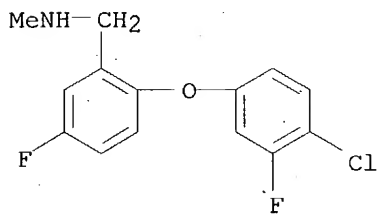
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-5-(trifluoromethyl)-
(9CI) (CA INDEX NAME)



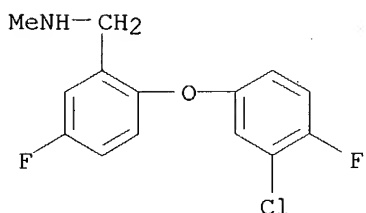
RN 289717-70-0 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4-methoxy-N-methyl- (9CI) (CA INDEX NAME)



RN 289717-71-1 HCAPLUS
 CN Benzenemethanamine, 2-(4-chloro-3-fluorophenoxy)-5-fluoro-N-methyl- (9CI) (CA INDEX NAME)



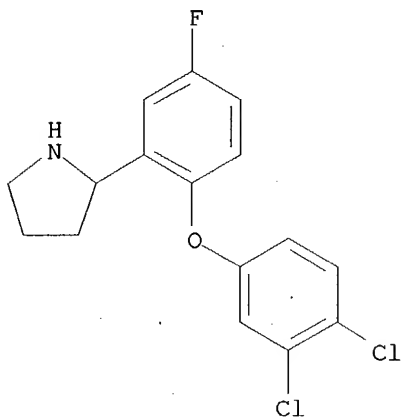
RN 289717-72-2 HCAPLUS
 CN Benzenemethanamine, 2-(3-chloro-4-fluorophenoxy)-5-fluoro-N-methyl- (9CI) (CA INDEX NAME)



RN 289717-73-3 HCPLUS

CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-, (-)- (9CI) (CA
INDEX NAME)

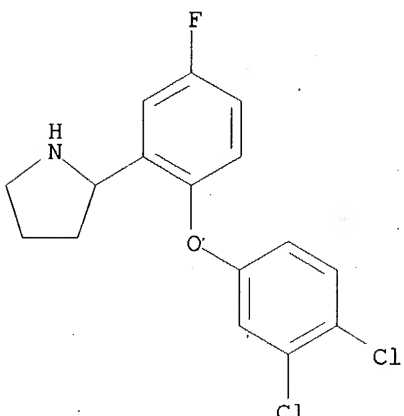
Rotation (-).



RN 289717-74-4 HCPLUS

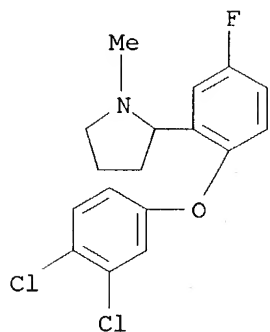
CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-, (+)- (9CI) (CA
INDEX NAME)

Rotation (+).



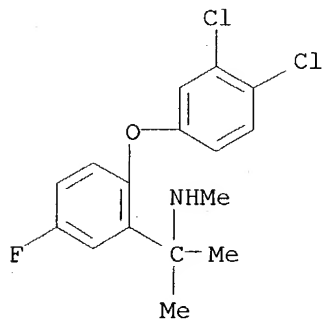
RN 289717-75-5 HCPLUS

CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-1-methyl- (9CI)
(CA INDEX NAME)



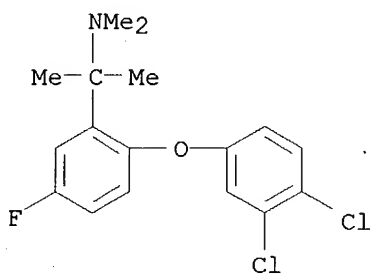
RN 444888-21-5 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N,α,α-trimethyl- (9CI) (CA INDEX NAME)



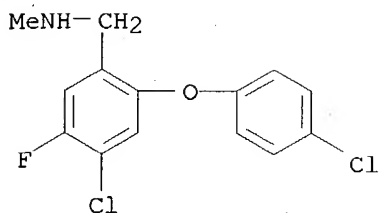
RN 444888-22-6 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N,N,α,α-tetramethyl- (9CI) (CA INDEX NAME)

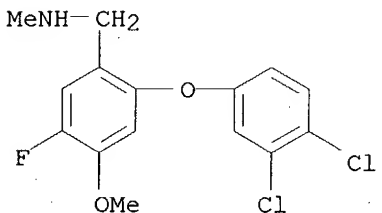


RN 444888-23-7 HCPLUS

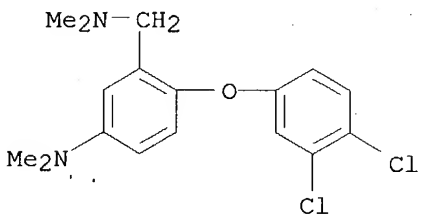
CN Benzenemethanamine, 4-chloro-2-(4-chlorophenoxy)-5-fluoro-N-methyl- (9CI) (CA INDEX NAME)



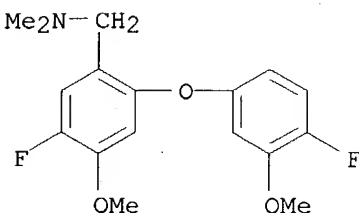
RN 444888-24-8 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-4-methoxy-N-methyl-
(9CI) (CA INDEX NAME)

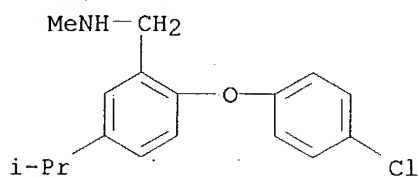
RN 444888-25-9 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-(dimethylamino)-N,N-dimethyl-
(9CI) (CA INDEX NAME)

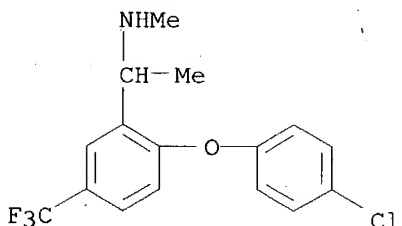
RN 444888-26-0 HCPLUS

CN Benzenemethanamine, 5-fluoro-2-(4-fluoro-3-methoxyphenoxy)-4-methoxy-N,N-dimethyl-
(9CI) (CA INDEX NAME)

RN 444888-27-1 HCAPLUS

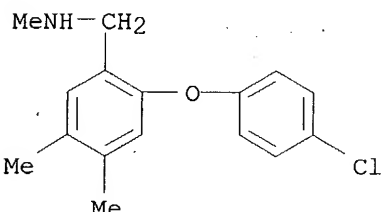
CN Benzenemethanamine, 2-(4-chlorophenoxy)-N-methyl-5-(1-methylethyl)- (9CI)
(CA INDEX NAME)

RN 444888-28-2 HCAPLUS

CN Benzenemethanamine, 2-(4-chlorophenoxy)-N, α -dimethyl-5-(trifluoromethyl)- (9CI) (CA INDEX NAME)

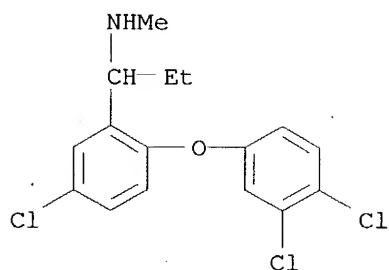
RN 444888-29-3 HCAPLUS

CN Benzenemethanamine, 2-(4-chlorophenoxy)-N,4,5-trimethyl- (9CI) (CA INDEX NAME)

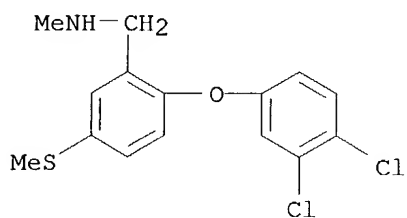


RN 444888-30-6 HCAPLUS

CN Benzenemethanamine, 5-chloro-2-(3,4-dichlorophenoxy)- α -ethyl-N-methyl- (9CI) (CA INDEX NAME)

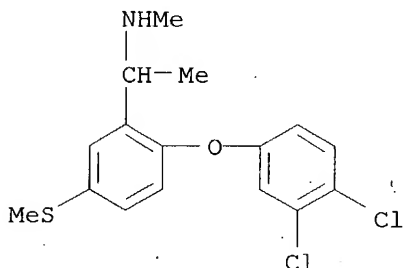


RN 444888-31-7 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-5-(methylthio)- (9CI)
(CA INDEX NAME)

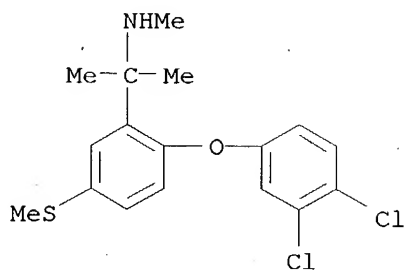
RN 444888-32-8 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,α-dimethyl-5-(methylthio)- (9CI) (CA INDEX NAME)



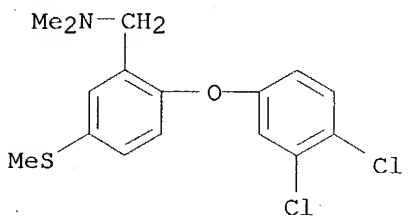
RN 444888-33-9 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,α,α-trimethyl-5-(methylthio)- (9CI) (CA INDEX NAME)



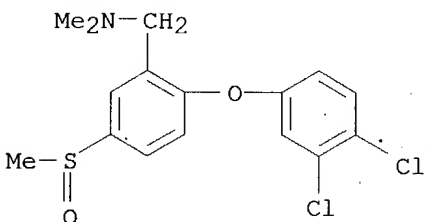
RN 444888-34-0 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-5-(methylthio)-(9CI) (CA INDEX NAME)



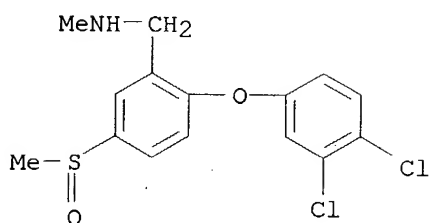
RN 444888-35-1 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-5-(methylsulfinyl)-(9CI) (CA INDEX NAME)

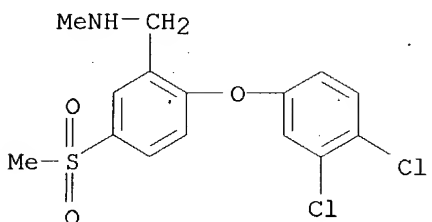


RN 444888-36-2 HCPLUS

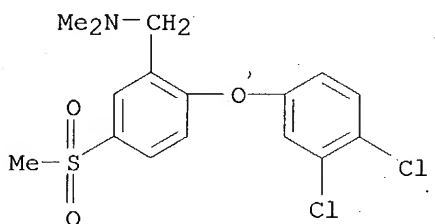
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-5-(methylsulfinyl)-(9CI) (CA INDEX NAME)



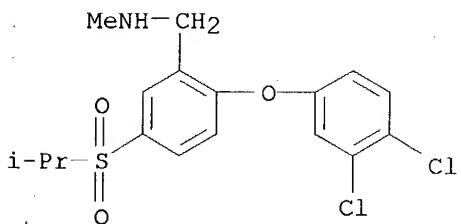
RN 444888-37-3 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-5-(methylsulfonyl)-(9CI) (CA INDEX NAME)



RN 444888-38-4 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-5-(methylsulfonyl)-(9CI) (CA INDEX NAME)

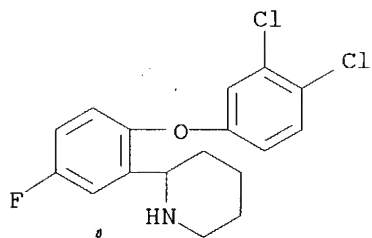


RN 444888-39-5 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-5-[(1-methylethyl)sulfonyl]-(9CI) (CA INDEX NAME)

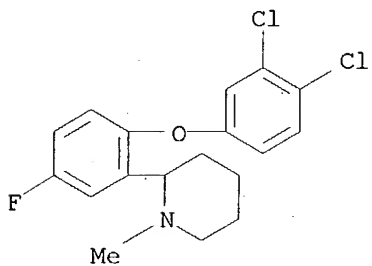


RN 444888-40-8 HCPLUS

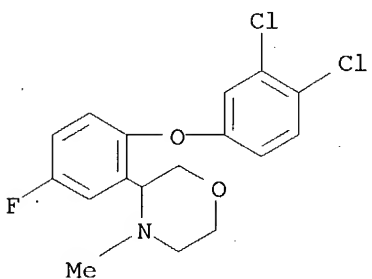
CN Piperidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]- (9CI) (CA INDEX NAME)



RN 444888-41-9 HCPLUS

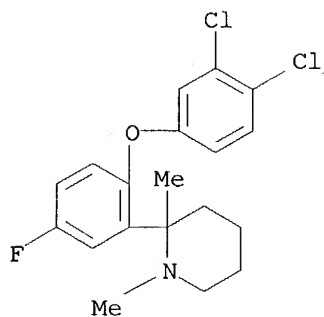
CN Piperidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-1-methyl- (9CI)
(CA INDEX NAME)

RN 444888-42-0 HCPLUS

CN Morpholine, 3-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-4-methyl- (9CI)
(CA INDEX NAME)

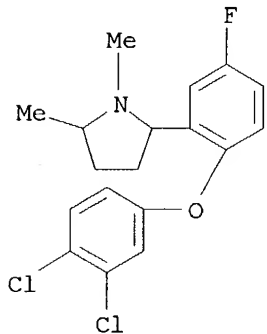
RN 444888-43-1 HCPLUS

CN Piperidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-1,2-dimethyl- (9CI)
(CA INDEX NAME)



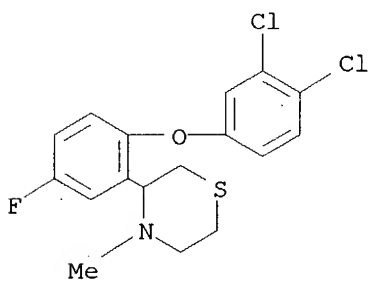
RN 444888-45-3 HCAPLUS

CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-1,5-dimethyl- (9CI) (CA INDEX NAME)



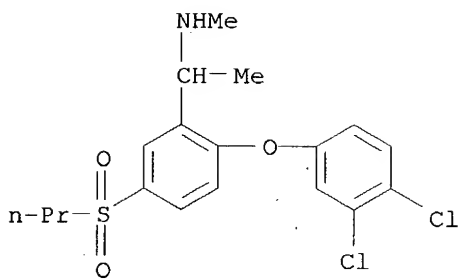
RN 444888-46-4 HCAPLUS

CN Thiomorpholine, 3-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-4-methyl- (9CI) (CA INDEX NAME)

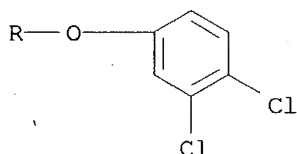
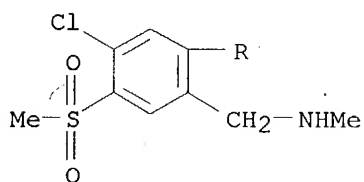


RN 444888-48-6 HCAPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,α-dimethyl-5-(propylsulfonyl)- (9CI) (CA INDEX NAME)

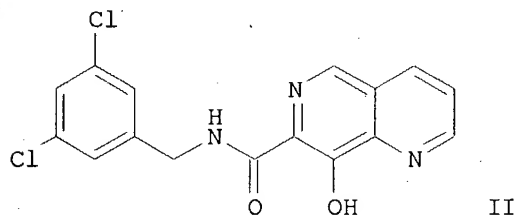
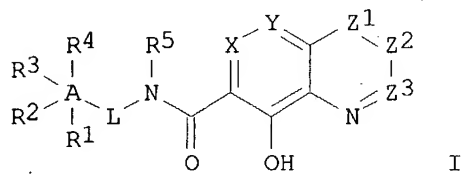


RN 444888-49-7 HCPLUS
 CN Benzenemethanamine, 4-chloro-2-(3,4-dichlorophenoxy)-N-methyl-5-(methylsulfonyl)- (9CI) (CA INDEX NAME)



L51 ANSWER 14 OF 44 HCPLUS COPYRIGHT 2004 ACS on STN
 ACCESSION NUMBER: 2002:293652 HCPLUS
 DOCUMENT NUMBER: 136:325531
 TITLE: Preparation of (poly)azanaphthalenyl carboxamides as HIV integrase inhibitors
 INVENTOR(S): Anthony, Neville J.; Gomez, Robert P.; Young, Steven D.; Egbertson, Melissa; Wai, John S.; Zhuang, Linghang; Embrey, Mark; Tran, Lekhanh; Melamed, Jeffrey Y.; Langford, H. Marie; Guare, James P.; Fisher, Thorsten E.; Jolly, Samson M.; Kuo, Michelle S.; Perlow, Debra S.; Bennett, Jennifer J.; Funk, Timothy W.
 PATENT ASSIGNEE(S): Merck & Co., Inc., USA
 SOURCE: PCT Int. Appl., 434 pp.
 CODEN: PIXXD2
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 2
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2002030930	A2	20020418	WO 2001-US31456	20011009
W:	AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM			
RW:	GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG			
AU 2002011527	A5	20020422	AU 2002-11527	20011009
EP 1326865	A2	20030716	EP 2001-979582	20011009
R:	AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR			
US 2003055071	A1	20030320	US 2001-973853	20011010
PRIORITY APPLN. INFO.:			US 2000-239707P	P 20001012
			US 2001-281656P	P 20010405
			WO 2001-US31456	W 20011009
OTHER SOURCE(S):	MARPAT 136:325531			
GI				



AB Title compds., including certain quinoline carboxamide and naphthyridine carboxamide derivs., I [wherein A = (un)substituted Ph or Ph fused to a carbocycle; L = a single bond, or (un)substituted alkyl, alkenyl, alkylcycloalkylalkyl, or alkyl-M-alkyl; M = NR_a, OCO, or CO₂; X = N or CQ₁; Y = N or CQ₂, provided that X and Y are not both N; Z₁ = N or CQ₃; Z₂ = N or CQ₄; Z₃ = N or CH; Q₁-Q₄ = independently H, halo, CN, NR₁CR₁O, or (un)substituted alkyl, alkoxy, alkenyl, alkynyl, carbamoyl, carboximidamido, amino, etc.; or C₂Q₂Q₃ = (un)substituted 5- or 6-membered carbocycle or heterocycle; R₁ and R₂ = independently H, OH, halo, NO₂, CN, or (un)substituted alkyl, alkenyl, alkoxy, amino, sulfonylamino, etc.; R₃ and R₄ = independently H, halo, CN, NO₂, OH, alkenyl, or (un)substituted alkyl, amino, sulfonylamino, etc.; R₅ = H, CN, CN, or (un)substituted

alkyl or aryl; Ra = independently H or (halo)alkyl; or pharmaceutically acceptable salts thereof] were prepared I are inhibitors of HIV integrase and inhibitors of HIV replication, and are useful in the prevention or treatment of infection by HIV and the treatment of AIDS, as compds. or pharmaceutically acceptable salts, or as ingredients in pharmaceutical compns., optionally in combination with other antivirals, immunomodulators, antibiotics, or vaccines. For example, Mitsunobu reaction of iso-Pr 3-(hydroxymethyl)pyridine-2-carboxylate with Me N-[(4-methylphenyl)sulfonyl]glycinate, followed by cyclization in the presence on NaOMe, afforded Me 8-hydroxy-1,6-naphthyridine-7-carboxylate. Coupling with 3,5-dichlorobenzylamine in toluene gave II. Representative compds. were assayed for the inhibition of acute HIV infection of T-lymphoid cells and demonstrated IC₅₀ values of < 20 μM.

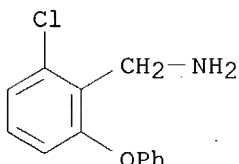
IT 175136-89-7, 2-Aminomethyl 3-chlorodiphenyl ether

RL: RCT (Reactant); RACT (Reactant or reagent)

(reactant; preparation of (poly)azanaphthalenyl carboxamides as HIV integrase inhibitors for treatment of AIDS)

RN 175136-89-7 HCPLUS

CN Benzenemethanamine, 2-chloro-6-phenoxy- (9CI) (CA INDEX NAME)



L51 ANSWER 15 OF 44 HCPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: 2002:171851 HCPLUS

DOCUMENT NUMBER: 136:232110

TITLE: Preparation of phenoxybenzylamines as selective serotonin re-uptake inhibitors

INVENTOR(S): Adam, Mavis Diane; Andrews, Mark David; Elliott, Mark Leonard; Gymer, Geoffrey Edward; Hepworth, David; Howard, Harry Ralph, Jr.; Middleton, Donald Stuart; Stobie, Alan

PATENT ASSIGNEE(S): Pfizer Limited, UK; Pfizer Inc.

SOURCE: PCT Int. Appl., 110 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

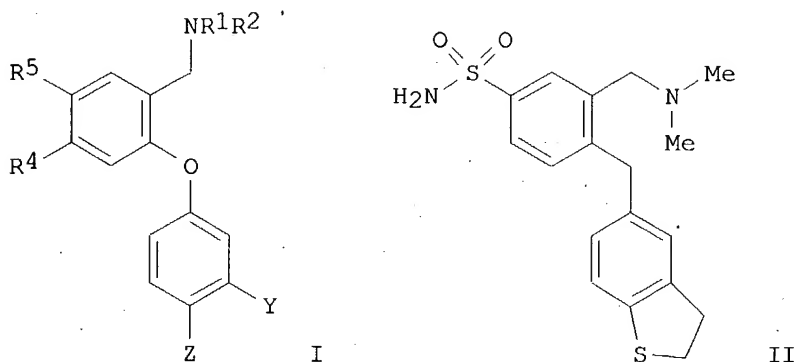
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2002018333	A1	20020307	WO 2001-IB1521	20010822
W:	AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM			

RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
BJ, CF, CG, CI, CM, GA, GN, GO, GW, ML, MR, NE, SN, TD, TG
AU 2001078650 A5 20020313 AU 2001-78650 20010822
EP 1313701 A1 20030528 EP 2001-956734 20010822
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
BR 2001013610 A 20030624 BR 2001-13610 20010822
US 2003060456 A1 20030327 US 2001-941177 20010827
US 6610747 B2 20030826
BG 107544 A 20031031 BG 2003-107544 20030207
NO 2003000842 A 20030428 NO 2003-842 20030224
HR 2003000141 A1 20030430 HR 2003-141 20030226
PRIORITY APPLN. INFO.: GB 2000-21593 A 20000831
GB 2001-7116 A 20010321
US 2000-240271P P 20001013
US 2001-292400P P 20010521
WO 2001-IB1521 W 20010822

OTHER SOURCE(S): MARPAT 136:232110

GI



AB Title compds. I [R1 and R2 independently = H, alkyl or (CH₂)_n(C₃-C₆cycloalkyl) wherein n = 0, 1, 2 or 3; or R1 and R2 together with the nitrogen to which they are attached from an azetidine ring; Z or Y is -SR₃ and the other Z or Y is halogen or -R₃; wherein R₃ = C₁₋₄ alkyl optionally substituted with fluorine; except that R₃ is not CF₃; or Z and Y are linked so that, together with the interconnecting atoms, Z and Y form a fused 5 to 7-membered carbocyclic or heterocyclic ring, and wherein when Z and Y form a heterocyclic ring, in addition to carbon atoms, the linkage contains one or two heteroatoms independently selected from O, S and N; R₄ and R₅ independently = A-X, wherein A = -CH=CH- or -(CH₂)_p- where p is 0, 1 or 2; X = H, halo, CONR₆R₇, SO₂NR₆R₇, SO₂NHC(=O)R₆, OH, C₁₋₄alkoxy, etc; or A-X = (un)substituted 5- or 6-membered heterocyclic ring containing 1, 2 or 3 heteroatoms selected from N, S and O; R₆ and R₇ independently = H, (un)substituted alkyl; or R₆ and R₇ together with the N to which they are attached form a (un)substituted 4-6 membered heterocyclic ring] and there pharmaceutically acceptable salts are prepared

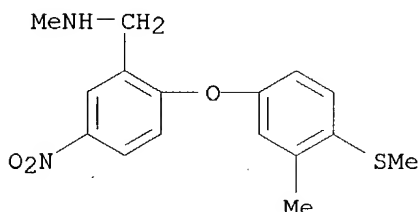
Thus, II was prepared via substitution of 5-(aminosulfonyl)-2-fluoro-N-methylbenzamide by 2,3-dihydrobenzo[b]thiophen-5-ol with successive $\text{BF}_3\cdot\text{THF}$ catalyzed amide reduction, formylation of secondary amine, and reduction. II demonstrated a serotonin re-uptake inhibition IC₅₀ of 4.7nM. I inhibit monoamine re-uptake and in particular exhibit activity as selective serotonin reuptake inhibitors.

IT 402912-23-6P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
(intermediate; preparation of serotonin re-uptake inhibitors
phenoxybenzylamines)

RN 402912-23-6 HCPLUS

CN Benzenemethanamine, N-methyl-2-[3-methyl-4-(methylthio)phenoxy]-5-nitro-(9CI) (CA INDEX NAME)



IT 402910-43-4P 402910-50-3P 402910-61-6P

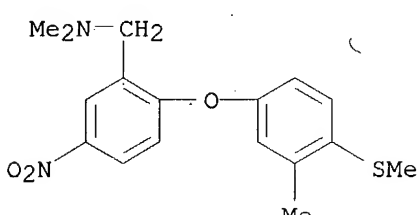
402910-62-7P 402910-90-1P 402910-96-7P

402911-10-8P 402911-12-0P 402911-20-0P

RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses)
(target compound; preparation of serotonin re-uptake inhibitors
phenoxybenzylamines)

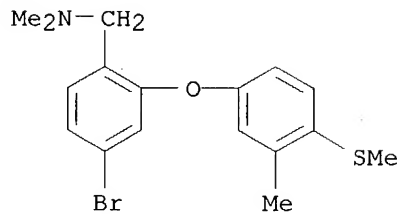
RN 402910-43-4 HCPLUS

CN Benzenemethanamine, N,N-dimethyl-2-[3-methyl-4-(methylthio)phenoxy]-5-nitro- (9CI) (CA INDEX NAME)

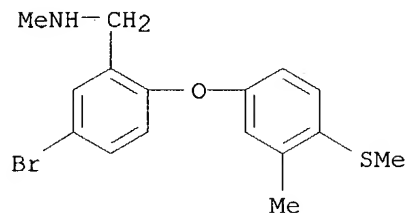


RN 402910-50-3 HCPLUS

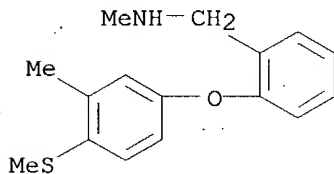
CN Benzenemethanamine, 4-bromo-N,N-dimethyl-2-[3-methyl-4-(methylthio)phenoxy]- (9CI) (CA INDEX NAME)



RN 402910-61-6 HCAPLUS

CN Benzenemethanamine, 5-bromo-N-methyl-2-[3-methyl-4-(methylthio)phenoxy]-
(9CI) (CA INDEX NAME)

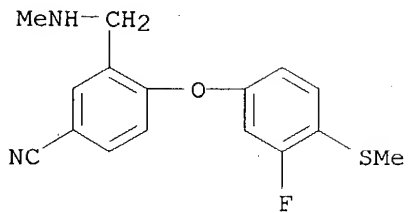
RN 402910-62-7 HCAPLUS

CN Benzenemethanamine, N-methyl-2-[3-methyl-4-(methylthio)phenoxy]-,
hydrochloride (9CI) (CA INDEX NAME)

● HCl

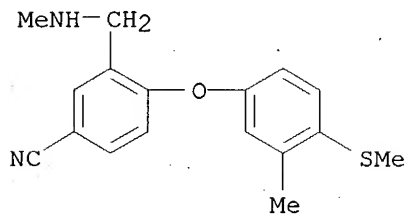
RN 402910-90-1 HCAPLUS

CN Benzonitrile, 4-[3-fluoro-4-(methylthio)phenoxy]-3-[(methylamino)methyl]-,
monohydrochloride (9CI) (CA INDEX NAME)



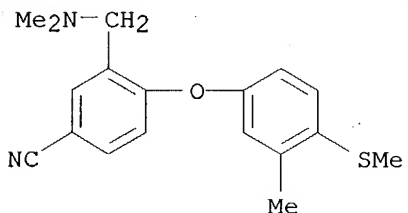
● HCl

RN 402910-96-7 HCPLUS
 CN Benzonitrile, 3-[(methylamino)methyl]-4-[3-methyl-4-(methylthio)phenoxy]-, monohydrochloride (9CI) (CA INDEX NAME)



● HCl

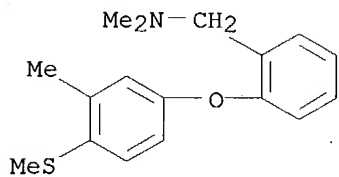
RN 402911-10-8 HCPLUS
 CN Benzonitrile, 3-[(dimethylamino)methyl]-4-[3-methyl-4-(methylthio)phenoxy]-, monohydrochloride (9CI) (CA INDEX NAME)



● HCl

RN 402911-12-0 HCPLUS
 CN Benzenemethanamine, N,N-dimethyl-2-[3-methyl-4-(methylthio)phenoxy]-,

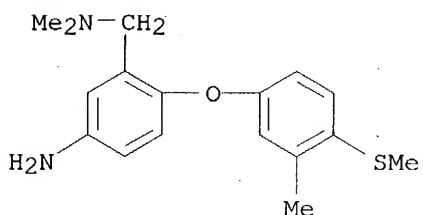
hydrochloride (9CI) (CA INDEX NAME)



● HCl

RN 402911-20-0 HCPLUS

CN Benzenemethanamine, 5-amino-N,N-dimethyl-2-[3-methyl-4-(methylthio)phenoxy]- (9CI) (CA INDEX NAME)



IT 402910-41-2P 402910-54-7P 402910-56-9P

402910-59-2P 402910-60-5P 402911-11-9P

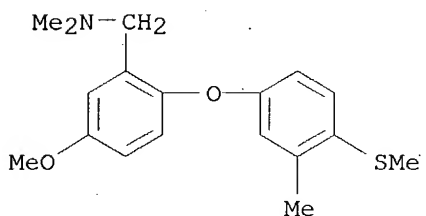
402911-49-3P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(target compound; preparation of serotonin re-uptake inhibitors
phenoxybenzylamines)

RN 402910-41-2 HCPLUS

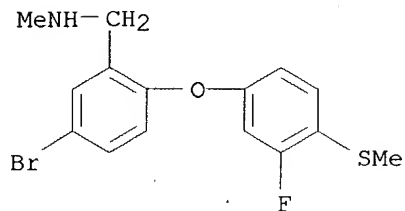
CN Benzenemethanamine, 5-methoxy-N,N-dimethyl-2-[3-methyl-4-(methylthio)phenoxy]-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

RN 402910-54-7 HCPLUS

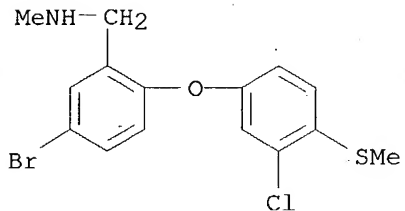
CN Benzenemethanamine, 5-bromo-2-[3-fluoro-4-(methylthio)phenoxy]-N-methyl-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

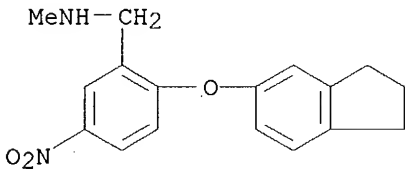
RN 402910-56-9 HCPLUS

CN Benzenemethanamine, 5-bromo-2-[3-chloro-4-(methylthio)phenoxy]-N-methyl- (9CI) (CA INDEX NAME)



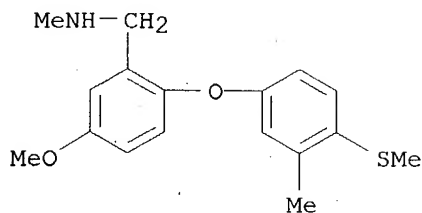
RN 402910-59-2 HCPLUS

CN Benzenemethanamine, 2-[(2,3-dihydro-1H-inden-5-yl)oxy]-N-methyl-5-nitro- (9CI) (CA INDEX NAME)



RN 402910-60-5 HCPLUS

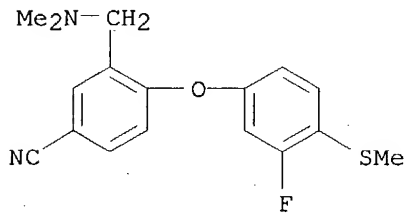
CN Benzenemethanamine, 5-methoxy-N-methyl-2-[3-methyl-4-(methylthio)phenoxy]-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

RN 402911-11-9 HCPLUS

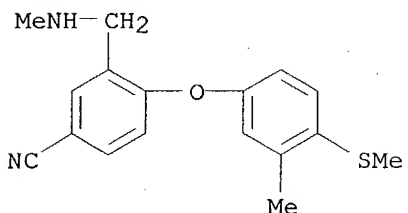
CN Benzonitrile, 3-[(dimethylamino)methyl]-4-[3-fluoro-4-(methylthio)phenoxy]-, monohydrochloride (9CI) (CA INDEX NAME)



● HCl

RN 402911-49-3 HCPLUS

CN Benzonitrile, 3-[(methylamino)methyl]-4-[3-methyl-4-(methylthio)phenoxy]- (9CI) (CA INDEX NAME)



REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L51 ANSWER 16 OF 44 HCPLUS COPYRIGHT 2004 ACS on STN

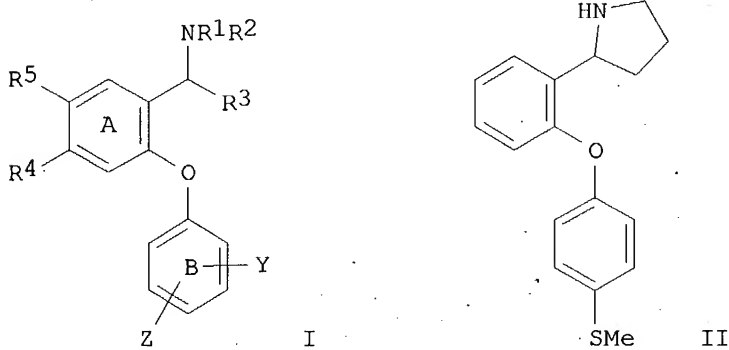
ACCESSION NUMBER: 2002:169117 HCPLUS

DOCUMENT NUMBER: 136:216641

TITLE: Preparation of phenoxyphenylheterocycles as selective serotonin reuptake inhibitors (SSRIs)
 INVENTOR(S): Andrews, Mark David; Hepworth, David; Middleton, Donald Stuart; Stobie, Alan
 PATENT ASSIGNEE(S): Pfizer Limited, UK; Pfizer Inc.
 SOURCE: Eur. Pat. Appl., 46 pp.
 CODEN: EPXXDW
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 1184372	A1	20020306	EP 2001-307032	20010817
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO				
US 2002183303	A1	20021205	US 2001-939475	20010824
US 6630504	B2	20031007		
BR 2001003797	A	20020604	BR 2001-3797	20010830
PRIORITY APPLN. INFO.:			GB 2000-21594	A 20000831
			GB 2001-5634	A 20010307
			US 2000-240326P	P 20001013

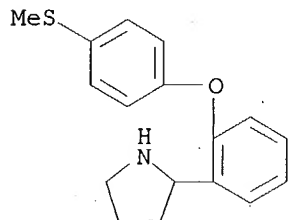
OTHER SOURCE(S): MARPAT 136:216641
GI



AB The title compds. [I; R1 = H, alkyl; R2, R3, together with the interconnecting atoms, form a 4-8 membered saturated ring containing 1-2 heteroatoms (including the N atom to which R2 is attached) wherein a second heteroatom, if present, is selected from O, N and S; Z = CF₃, OCF₃, alkylthio or alkoxy; Y = H, halo, ORa, Ra or alkylthio (wherein Ra = alkyl optionally substituted with F atoms); or when Z and Y are attached para and meta to the ether linkage linking rings A and B, Z and Y are linked so that, together with the interconnecting atoms, Z and Y form a fused 5-7 membered carbocyclic or heterocyclic ring which may be saturated, unsatd. or aromatic; R4, R5 = AX (wherein A = CH:CH, (CH₂)_p; p = 0-2; X = H, F, Cl, OH, etc.)] and their salts, useful for the treatment of prevention of a disorder in which the regulation of monoamine transporter function is implicated, were prepared and formulated. Thus, reducing

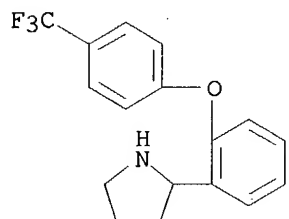
5-[2-[4-(methylsulfanyl)phenoxy]phenyl]-3,4-dihydro-2H-pyrrole (preparation given) with NaBH₄ in EtOH afforded 72% II.HCl which showed IC₅₀ of ≤ 100 nM in test for serotonin re-uptake inhibition.

IT 402714-37-8P 402714-38-9P 402714-39-0P
 402714-48-1P 402714-52-7P 402714-53-8P
 RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses)
 (preparation of phenoxyphenylpyrrolidines as selective serotonin reuptake inhibitors (SSRIs))
 RN 402714-37-8 HCAPLUS
 CN Pyrrolidine, 2-[2-[4-(methylthio)phenoxy]phenyl]-, hydrochloride (9CI)
 (CA INDEX NAME)



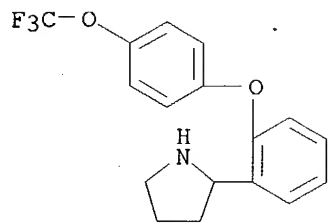
● HCl

RN 402714-38-9 HCAPLUS
 CN Pyrrolidine, 2-[2-[4-(trifluoromethyl)phenoxy]phenyl]-, hydrochloride (9CI) (CA INDEX NAME)



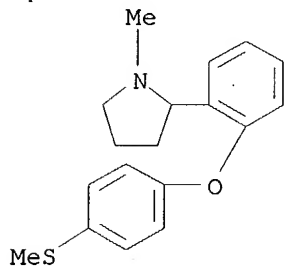
● HCl

RN 402714-39-0 HCAPLUS
 CN Pyrrolidine, 2-[2-[4-(trifluoromethoxy)phenoxy]phenyl]-, hydrochloride (9CI) (CA INDEX NAME)



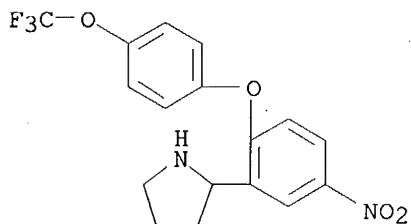
● HCl

RN 402714-48-1 HCPLUS
 CN Pyrrolidine, 1-methyl-2-[2-[4-(methylthio)phenoxy]phenyl]-, hydrochloride
 (9CI) (CA INDEX NAME)

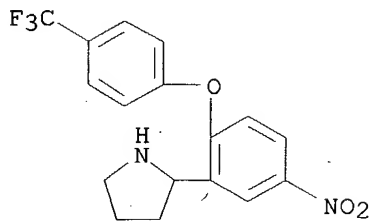


● HCl

RN 402714-52-7 HCPLUS
 CN Pyrrolidine, 2-[5-nitro-2-[4-(trifluoromethoxy)phenoxy]phenyl]- (9CI) (CA
 INDEX NAME)



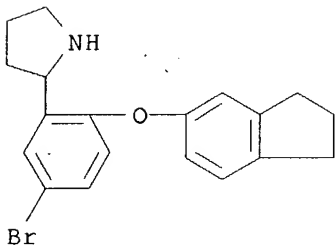
RN 402714-53-8 HCPLUS
 CN Pyrrolidine, 2-[5-nitro-2-[4-(trifluoromethyl)phenoxy]phenyl]- (9CI) (CA
 INDEX NAME)



IT 402714-40-3P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
 (preparation of phenoxyphenylpyrrolidines as selective serotonin reuptake inhibitors (SSRIs))

RN 402714-40-3 HCAPLUS

CN Pyrrolidine, 2-[5-bromo-2-[(2,3-dihydro-1H-inden-5-yl)oxy]phenyl]- (9CI)
 (CA INDEX NAME)

IT 402714-74-3P 402714-75-4P 402714-94-7P

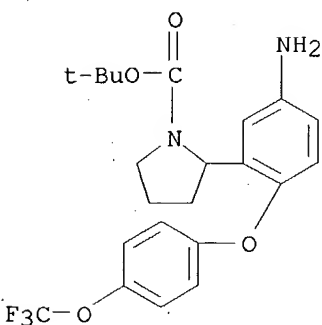
402714-95-8P 402714-96-9P 402714-97-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of phenoxyphenylpyrrolidines as selective serotonin reuptake inhibitors (SSRIs))

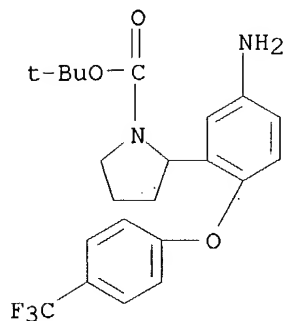
RN 402714-74-3 HCAPLUS

CN 1-Pyrrolidinecarboxylic acid, 2-[5-amino-2-[4-(trifluoromethoxy)phenoxy]phenyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)



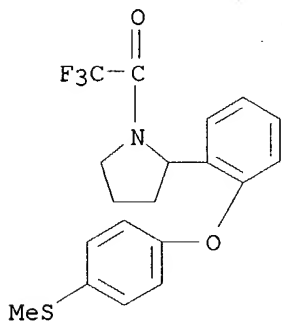
RN 402714-75-4 HCAPLUS

CN 1-Pyrrolidinecarboxylic acid, 2-[5-amino-2-[4-(trifluoromethyl)phenoxy]phenyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)



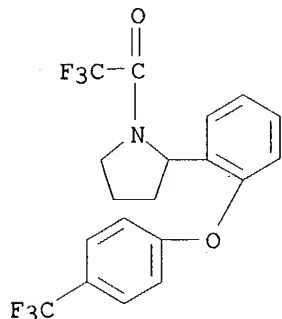
RN 402714-94-7 HCAPLUS

CN Pyrrolidine, 2-[2-[4-(methylthio)phenoxy]phenyl]-1-(trifluoroacetyl)-(9CI) (CA INDEX NAME)



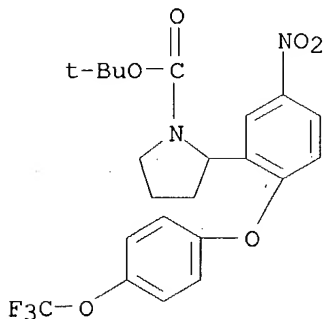
RN 402714-95-8 HCAPLUS

CN Pyrrolidine, 1-(trifluoroacetyl)-2-[2-[4-(trifluoromethyl)phenoxy]phenyl]-(9CI) (CA INDEX NAME)



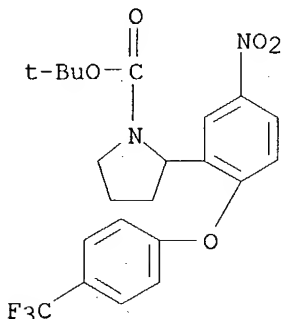
RN 402714-96-9 HCPLUS

CN 1-Pyrrolidinecarboxylic acid, 2-[5-nitro-2-[4-(trifluoromethoxy)phenoxy]phenyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)



RN 402714-97-0 HCPLUS

CN 1-Pyrrolidinecarboxylic acid, 2-[5-nitro-2-[4-(trifluoromethyl)phenoxy]phenyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)



REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L51 ANSWER 17 OF 44 HCPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: 2001:730683 HCPLUS

DOCUMENT NUMBER: 135:288572

TITLE: Preparation of diphenyl ether compounds as serotonin re-uptake inhibitors

INVENTOR(S): Andrews, Mark David; Hepworth, David; Middleton, Donald Stuart; Stobie, Alan

PATENT ASSIGNEE(S): Pfizer Limited, UK; Pfizer Inc.

SOURCE: PCT Int. Appl., 158 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

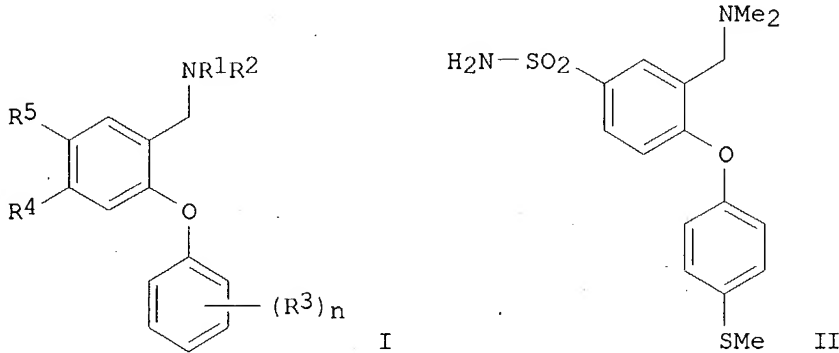
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2001072687	A1	20011004	WO 2001-IB428	20010319
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM				
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG				
US 2002052395	A1	20020502	US 2001-810378	20010316
US 6448293	B2	20020910		
EP 1268396	A1	20030102	EP 2001-917347	20010319
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR				
BR 2001009547	A	20030610	BR 2001-9547	20010319
NZ 519972	A	20030725	NZ 2001-519972	20010319
JP 2003528845	T2	20030930	JP 2001-570602	20010319
BG 106912	A	20030131	BG 2002-106912	20020709
NO 2002004663	A	20020927	NO 2002-4663	20020927
PRIORITY APPLN. INFO.:			GB 2000-7884	A 20000331
			US 2000-197127P	P 20000414
			WO 2001-IB428	W 20010319

OTHER SOURCE(S):

MARPAT 135:288572

GI

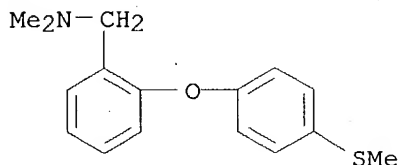


AB Title compds. I [wherein R₁ and R₂ = independently H or (cycloalkyl)alkyl; or R₁ and R₂ together with the N to which they are attached form an azetidine ring; R₃ = independently CF₃, OC₂F₅, alkylthio, or alkoxy; n = 1-3; R₄ and R₅ = independently AX; A = CH:CH or (CH₂)_p; p = 0-2; X = H, halo, OH, alkoxy, NO₂, CN, CHO, alkylthio, alkylsulfinyl, alkylsulfonyl, or (un)substituted carbamoyl, sulfamoyl, amino, carboxy, etc.; or pharmaceutically acceptable salts, solvates, or polymorphs thereof] were prepared as monoamine re-uptake inhibitors, particularly as selective serotonin re-uptake inhibitors. For example, 4-(methylmercapto)phenol was

coupled with 2-fluorobenzaldehyde using K₂CO₃ in DMF to give 2-[4-(methylsulfanyl)phenoxy]benzaldehyde (100%). The aldehyde was dissolved in THF, DCM, Me₂NH•HCl, and TEA, treated with NaBH(OAc)₃, and converted to the salt with 1M HCl in Et₂O to afford N,N-dimethyl-N-[2-[4-(methylsulfanyl)phenoxy]benzyl]amine•HCl (84%). Coupling the salt with ClSO₃H in CH₂Cl₂ at 0° to 5°C, followed by stepwise addition of MeCN with POCl₃ and ammonia, produced the desired sulfonamide (II) in 61% yield. The latter showed serotonin re-uptake inhibition (SRI) activity with IC₅₀ ≤ 50 nM and was > 100-fold as potent in the inhibition of serotonin re-uptake than in the inhibition of dopamine and noradrenaline re-uptake. It is useful in the treatment of disorders such as depression, attention deficit hyperactivity disorder, obsessive-compulsive disorder, post-traumatic stress disorder, substance abuse disorders, and sexual dysfunction, including premature ejaculation (no data).

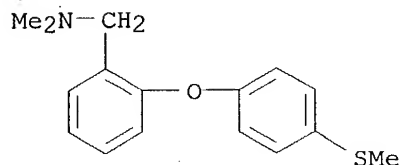
IT 289717-37-9P 364323-82-0P 364323-83-1P
 364323-84-2P 364323-85-3P 364323-86-4P
 364323-87-5P 364323-88-6P 364323-89-7P
 RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
 (intermediate; preparation of di-Ph ether compds. as serotonin re-uptake inhibitors)

RN 289717-37-9 HCPLUS
 CN Benzenemethanamine, N,N-dimethyl-2-[4-(methylthio)phenoxy]-, hydrochloride (9CI) (CA INDEX NAME)

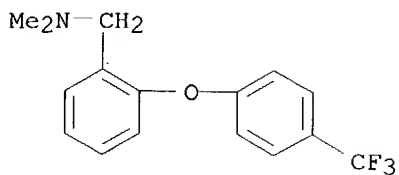


● HCl

RN 364323-82-0 HCPLUS
 CN Benzenemethanamine, N,N-dimethyl-2-[4-(methylthio)phenoxy]- (9CI) (CA INDEX NAME)



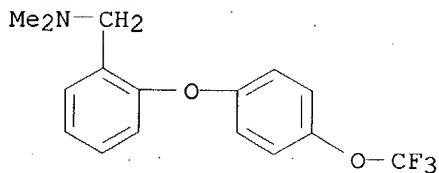
RN 364323-83-1 HCPLUS
 CN Benzenemethanamine, N,N-dimethyl-2-[4-(trifluoromethyl)phenoxy]-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

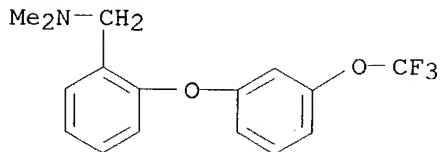
RN 364323-84-2 HCPLUS

CN Benzenemethanamine, N,N-dimethyl-2-[4-(trifluoromethoxy)phenoxy]- (9CI)
(CA INDEX NAME)



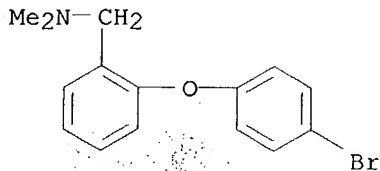
RN 364323-85-3 HCPLUS

CN Benzenemethanamine, N,N-dimethyl-2-[3-(trifluoromethoxy)phenoxy]- (9CI)
(CA INDEX NAME)



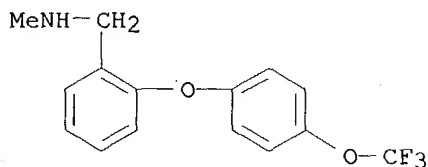
RN 364323-86-4 HCPLUS

CN Benzenemethanamine, 2-(4-bromophenoxy)-N,N-dimethyl- (9CI) (CA INDEX
NAME)

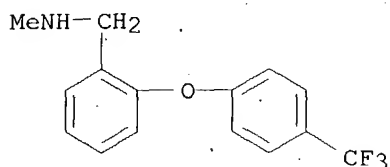


RN 364323-87-5 HCPLUS

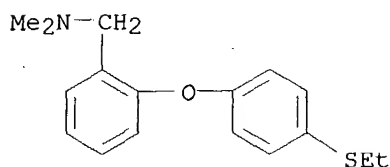
CN Benzenemethanamine, N-methyl-2-[4-(trifluoromethoxy)phenoxy]- (9CI) (CA
INDEX NAME)



RN 364323-88-6 HCPLUS
CN Benzenemethanamine, N-methyl-2-[4-(trifluoromethyl)phenoxy]- (9CI) (CA
INDEX NAME)



RN 364323-89-7 HCPLUS
CN Benzenemethanamine, 2-[4-(ethylthio)phenoxy]-N,N-dimethyl- (9CI) (CA
INDEX NAME)



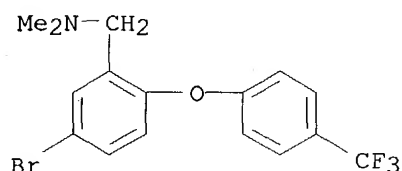
IT 364321-43-7P 364321-47-1P 364321-48-2P
364321-49-3P 364321-52-8P 364321-53-9P
364321-54-0P 364321-56-2P 364321-57-3P
364321-58-4P 364321-59-5P 364321-61-9P
364321-62-0P 364321-64-2P 364321-65-3P
364321-66-4P 364321-67-5P 364321-68-6P
364321-70-0P 364322-28-1P 364322-29-2P
364322-33-8P 364322-34-9P 364322-35-0P
364322-36-1P 364322-37-2P 364322-59-8P
364322-60-1P 364322-61-2P 364322-95-2P
364322-96-3P 364322-97-4P 364322-98-5P
364323-48-8P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses)

(preparation of di-Ph ether compds. as serotonin re-uptake inhibitors)

RN 364321-43-7 HCPLUS

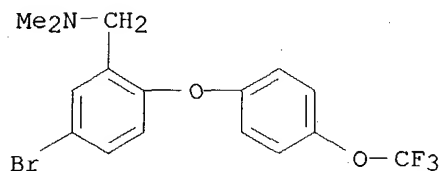
CN Benzenemethanamine, 5-bromo-N,N-dimethyl-2-[4-(trifluoromethyl)phenoxy]-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

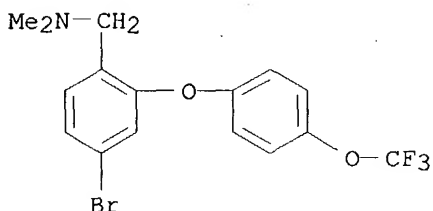
RN 364321-47-1 HCPLUS

CN Benzenemethanamine, 5-bromo-N,N-dimethyl-2-[4-(trifluoromethoxy)phenoxy]- (9CI) (CA INDEX NAME)



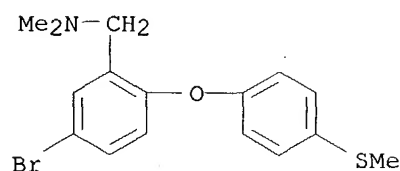
RN 364321-48-2 HCPLUS

CN Benzenemethanamine, 4-bromo-N,N-dimethyl-2-[4-(trifluoromethoxy)phenoxy]- (9CI) (CA INDEX NAME)



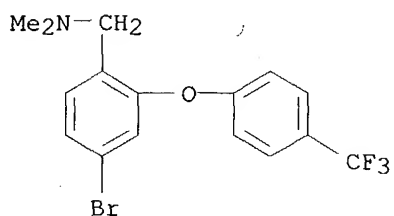
RN 364321-49-3 HCPLUS

CN Benzenemethanamine, 5-bromo-N,N-dimethyl-2-[4-(methylthio)phenoxy]-, hydrochloride (9CI) (CA INDEX NAME)

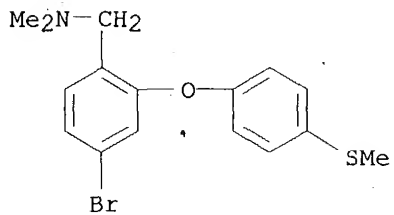


● HCl

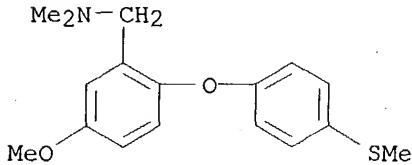
RN 364321-52-8 HCAPLUS

CN Benzenemethanamine, 4-bromo-N,N-dimethyl-2-[4-(trifluoromethyl)phenoxy]-
(9CI) (CA INDEX NAME)

RN 364321-53-9 HCAPLUS

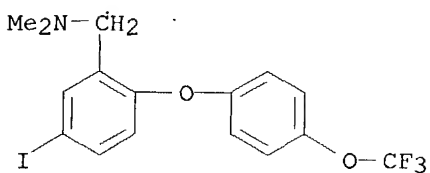
CN Benzenemethanamine, 4-bromo-N,N-dimethyl-2-[4-(methylthio)phenoxy]- (9CI)
(CA INDEX NAME)

RN 364321-54-0 HCAPLUS

CN Benzenemethanamine, 5-methoxy-N,N-dimethyl-2-[4-(methylthio)phenoxy]-
(9CI) (CA INDEX NAME)

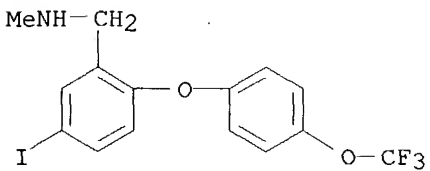
RN 364321-56-2 HCPLUS

CN Benzenemethanamine, 5-iodo-N,N-dimethyl-2-[4-(trifluoromethoxy)phenoxy]-(9CI) (CA INDEX NAME)



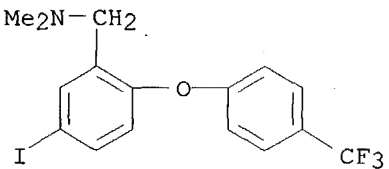
RN 364321-57-3 HCPLUS

CN Benzenemethanamine, 5-iodo-N-methyl-2-[4-(trifluoromethoxy)phenoxy]-(9CI) (CA INDEX NAME)



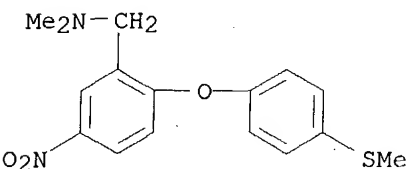
RN 364321-58-4 HCPLUS

CN Benzenemethanamine, 5-iodo-N,N-dimethyl-2-[4-(trifluoromethyl)phenoxy]-(9CI) (CA INDEX NAME)

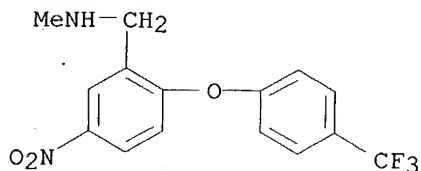


RN 364321-59-5 HCPLUS

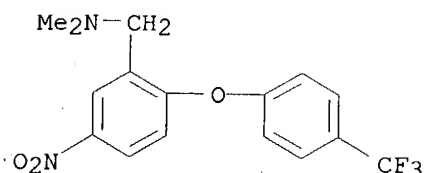
CN Benzenemethanamine, N,N-dimethyl-2-[4-(methylthio)phenoxy]-5-nitro-(9CI) (CA INDEX NAME)



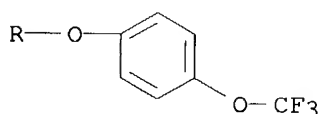
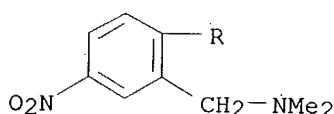
RN 364321-61-9 HCPLUS
 CN Benzenemethanamine, N-methyl-5-nitro-2-[4-(trifluoromethyl)phenoxy]- (9CI)
 (CA INDEX NAME)



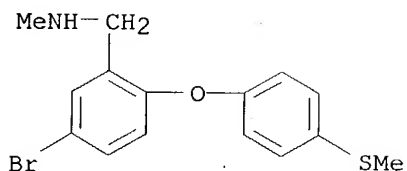
RN 364321-62-0 HCPLUS
 CN Benzenemethanamine, N,N-dimethyl-5-nitro-2-[4-(trifluoromethyl)phenoxy]- (9CI) (CA INDEX NAME)



RN 364321-64-2 HCPLUS
 CN Benzenemethanamine, N,N-dimethyl-5-nitro-2-[4-(trifluoromethoxy)phenoxy]- (9CI) (CA INDEX NAME)

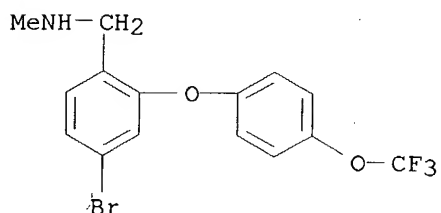


RN 364321-65-3 HCPLUS
 CN Benzenemethanamine, 5-bromo-N-methyl-2-[4-(methylthio)phenoxy]- (9CI) (CA INDEX NAME)



RN 364321-66-4 HCPLUS

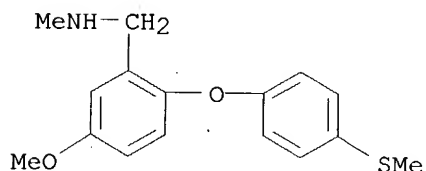
CN Benzenemethanamine, 4-bromo-N-methyl-2-[4-(trifluoromethoxy)phenoxy]-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

RN 364321-67-5 HCPLUS

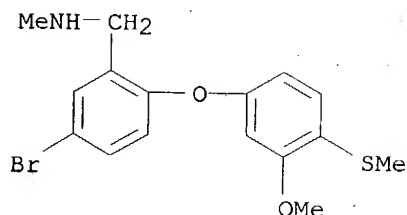
CN Benzenemethanamine, 5-methoxy-N-methyl-2-[4-(methylthio)phenoxy]-, hydrochloride (9CI) (CA INDEX NAME)



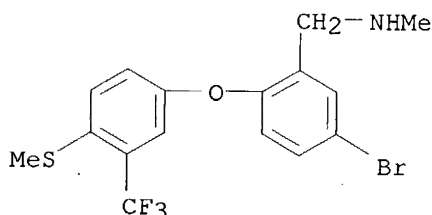
● HCl

RN 364321-68-6 HCPLUS

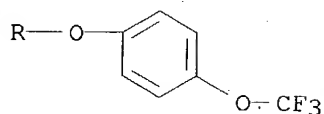
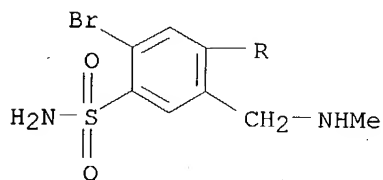
CN Benzenemethanamine, 5-bromo-2-[3-methoxy-4-(methylthio)phenoxy]-N-methyl-, (9CI) (CA INDEX NAME)



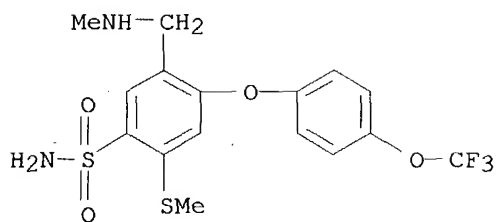
RN 364321-70-0 HCPLUS
 CN Benzenemethanamine, 5-bromo-*N*-methyl-2-[4-(methylthio)-3-(trifluoromethyl)phenoxy]- (9CI) (CA INDEX NAME)



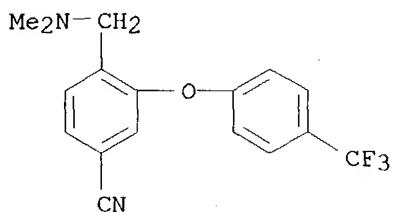
RN 364322-28-1 HCPLUS
 CN Benzenesulfonamide, 2-bromo-5-[(methylamino)methyl]-4-[4-(trifluoromethoxy)phenoxy]- (9CI) (CA INDEX NAME)



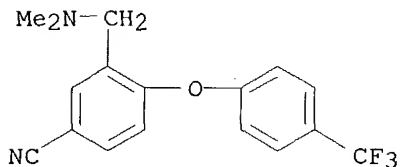
RN 364322-29-2 HCPLUS
 CN Benzenesulfonamide, 5-[(methylamino)methyl]-2-(methylthio)-4-[4-(trifluoromethoxy)phenoxy]- (9CI) (CA INDEX NAME)



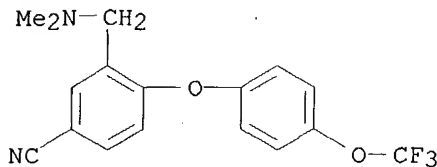
RN 364322-33-8 HCPLUS
 CN Benzonitrile, 4-[(dimethylamino)methyl]-3-[4-(trifluoromethyl)phenoxy]-
 (9CI) (CA INDEX NAME)



RN 364322-34-9 HCPLUS
 CN Benzonitrile, 3-[(dimethylamino)methyl]-4-[4-(trifluoromethyl)phenoxy]-
 (9CI) (CA INDEX NAME)

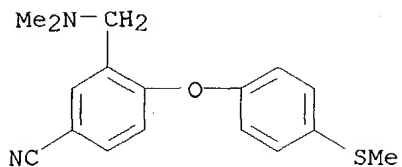


RN 364322-35-0 HCPLUS
 CN Benzonitrile, 3-[(dimethylamino)methyl]-4-[4-(trifluoromethoxy)phenoxy]-
 (9CI) (CA INDEX NAME)

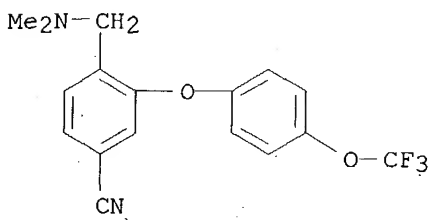


RN 364322-36-1 HCPLUS
 CN Benzonitrile, 3-[(dimethylamino)methyl]-4-[4-(methylthio)phenoxy]- (9CI)

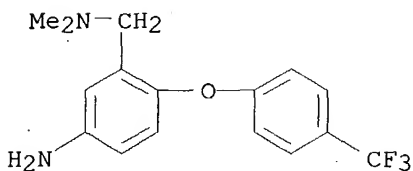
(CA INDEX NAME)



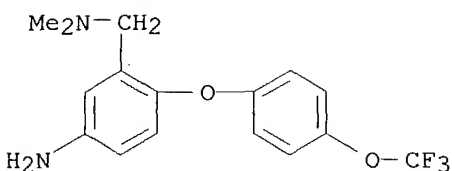
RN 364322-37-2 HCPLUS

CN Benzonitrile, 4-[(dimethylamino)methyl]-3-[4-(trifluoromethoxy)phenoxy]-
(9CI) (CA INDEX NAME)

RN 364322-59-8 HCPLUS

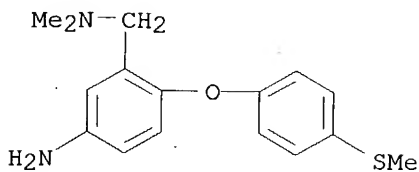
CN Benzenemethanamine, 5-amino-N,N-dimethyl-2-[4-(trifluoromethyl)phenoxy]-
(9CI) (CA INDEX NAME)

RN 364322-60-1 HCPLUS

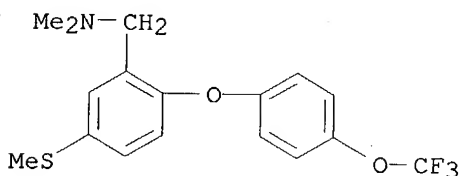
CN Benzenemethanamine, 5-amino-N,N-dimethyl-2-[4-(trifluoromethoxy)phenoxy]-
(9CI) (CA INDEX NAME)

RN 364322-61-2 HCPLUS

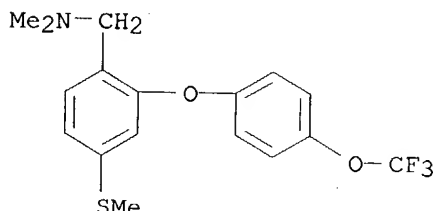
CN Benzenemethanamine, 5-amino-N,N-dimethyl-2-[4-(methylthio)phenoxy]- (9CI)
 (CA INDEX NAME)



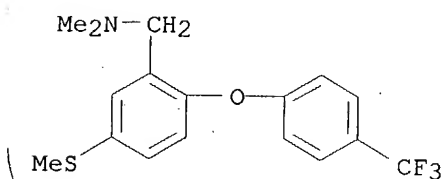
RN 364322-95-2 HCPLUS
 CN Benzenemethanamine, N,N-dimethyl-5-(methylthio)-2-[4-(trifluoromethoxy)phenoxy]- (9CI) (CA INDEX NAME)



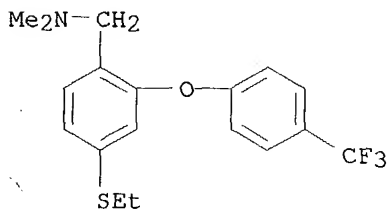
RN 364322-96-3 HCPLUS
 CN Benzenemethanamine, N,N-dimethyl-4-(methylthio)-2-[4-(trifluoromethoxy)phenoxy]- (9CI) (CA INDEX NAME)



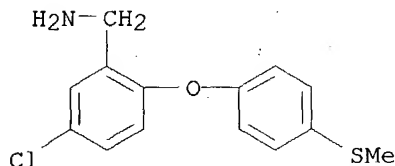
RN 364322-97-4 HCPLUS
 CN Benzenemethanamine, N,N-dimethyl-5-(methylthio)-2-[4-(trifluoromethyl)phenoxy]- (9CI) (CA INDEX NAME)



RN 364322-98-5 HCPLUS
 CN Benzenemethanamine, 4-(ethylthio)-N,N-dimethyl-2-[4-(trifluoromethyl)phenoxy]- (9CI) (CA INDEX NAME)



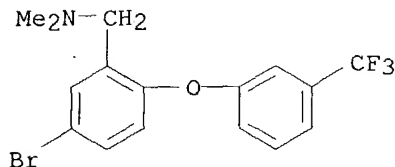
RN 364323-48-8 HCPLUS
 CN Benzenemethanamine, 5-chloro-2-[4-(methylthio)phenoxy]- (9CI) (CA INDEX NAME)



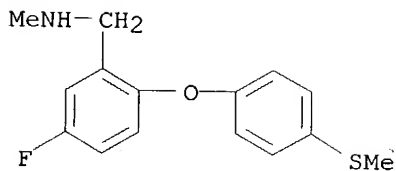
IT 364321-41-5P 364321-44-8P 364321-45-9P
 364321-50-6P 364321-51-7P 364321-55-1P
 364322-30-5P 364322-31-6P 364322-32-7P
 364322-38-3P 364322-89-4P 364322-99-6P
 364323-00-2P 364323-01-3P 364323-02-4P
364323-04-6P 364323-05-7P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
 (preparation of di-Ph ether compds. as serotonin re-uptake inhibitors)

RN 364321-41-5 HCPLUS
 CN Benzenemethanamine, 5-bromo-N,N-dimethyl-2-[3-(trifluoromethyl)phenoxy]- (9CI) (CA INDEX NAME)



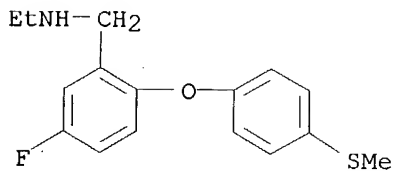
RN 364321-44-8 HCPLUS
 CN Benzenemethanamine, 5-fluoro-N-methyl-2-[4-(methylthio)phenoxy]-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

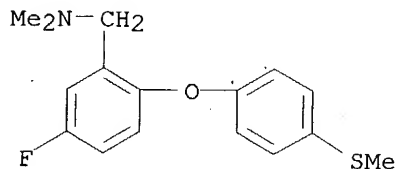
RN 364321-45-9 HCPLUS

CN Benzenemethanamine, N-ethyl-5-fluoro-2-[4-(methylthio)phenoxy]- (9CI) (CA INDEX NAME)



RN 364321-50-6 HCPLUS

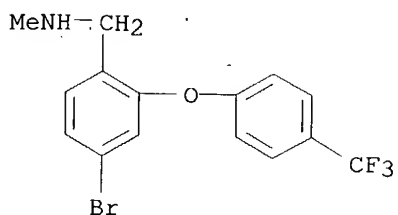
CN Benzenemethanamine, 5-fluoro-N,N-dimethyl-2-[4-(methylthio)phenoxy]-, hydrochloride (9CI) (CA INDEX NAME)



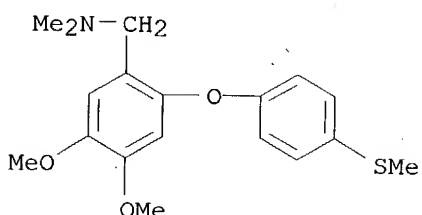
● HCl

RN 364321-51-7 HCPLUS

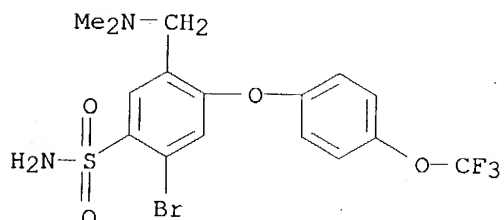
CN Benzenemethanamine, 4-bromo-N-methyl-2-[4-(trifluoromethyl)phenoxy]- (9CI) (CA INDEX NAME)



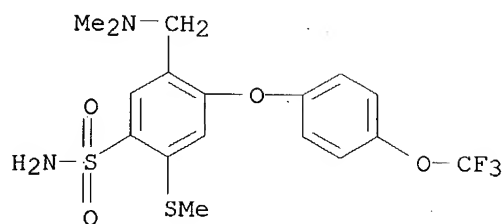
RN 364321-55-1 HCAPLUS
 CN Benzenemethanamine, 4,5-dimethoxy-N,N-dimethyl-2-[4-(methylthio)phenoxy]-
 (9CI) (CA INDEX NAME)



RN 364322-30-5 HCAPLUS
 CN Benzenesulfonamide, 2-bromo-5-[(dimethylamino)methyl]-4-[4-(trifluoromethoxy)phenoxy]- (9CI) (CA INDEX NAME)

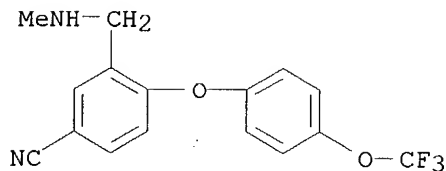


RN 364322-31-6 HCAPLUS
 CN Benzenesulfonamide, 5-[(dimethylamino)methyl]-2-(methylthio)-4-[4-(trifluoromethoxy)phenoxy]-, monohydrochloride (9CI) (CA INDEX NAME)

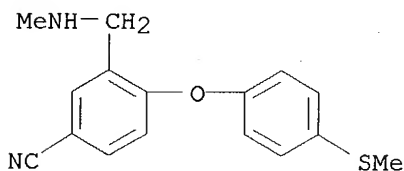


● HCl

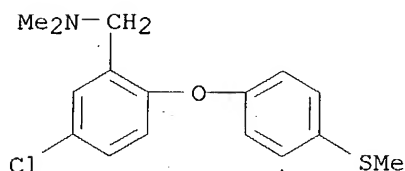
RN 364322-32-7 HCPLUS
 CN Benzonitrile, 3-[(methylamino)methyl]-4-[4-(trifluoromethoxy)phenoxy]-
 (9CI) (CA INDEX NAME)



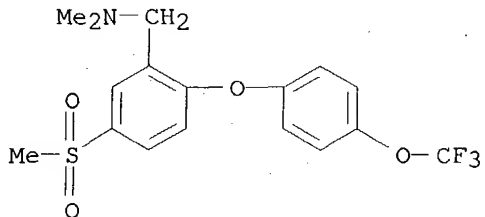
RN 364322-38-3 HCPLUS
 CN Benzonitrile, 3-[(methylamino)methyl]-4-[4-(methylthio)phenoxy]- (9CI)
 (CA INDEX NAME)



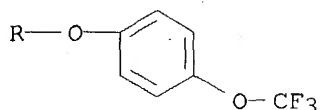
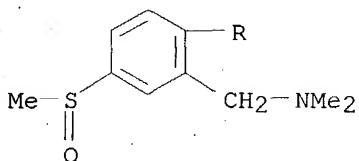
RN 364322-89-4 HCPLUS
 CN Benzenemethanamine, 5-chloro-N,N-dimethyl-2-[4-(methylthio)phenoxy]- (9CI)
 (CA INDEX NAME)



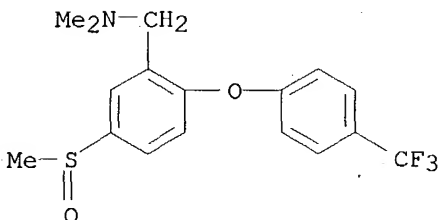
RN 364322-99-6 HCPLUS
 CN Benzenemethanamine, N,N-dimethyl-5-(methylsulfonyl)-2-[4-(trifluoromethoxy)phenoxy]- (9CI) (CA INDEX NAME)



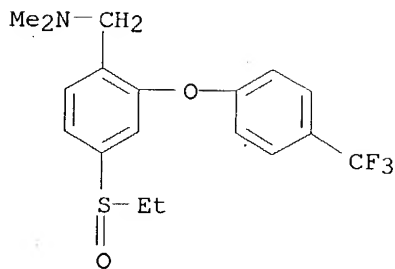
RN 364323-00-2 HCPLUS
 CN Benzenemethanamine, N,N-dimethyl-5-(methylsulfinyl)-2-[4-(trifluoromethoxy)phenoxy]- (9CI) (CA INDEX NAME)



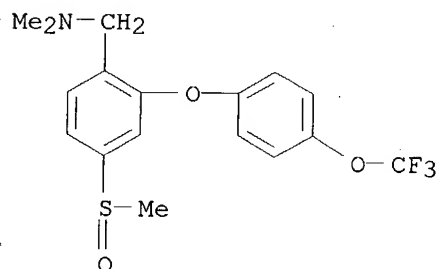
RN 364323-01-3 HCPLUS
 CN Benzenemethanamine, N,N-dimethyl-5-(methylsulfinyl)-2-[4-(trifluoromethyl)phenoxy]- (9CI) (CA INDEX NAME)



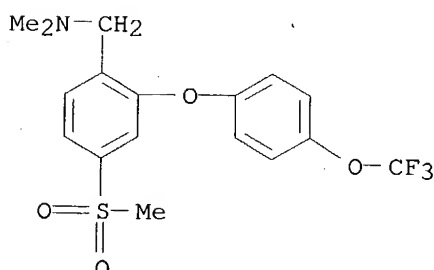
RN 364323-02-4 HCPLUS
 CN Benzenemethanamine, 4-(ethylsulfinyl)-N,N-dimethyl-2-[4-(trifluoromethyl)phenoxy]- (9CI) (CA INDEX NAME)



RN 364323-04-6 HCAPLUS
 CN Benzenemethanamine, N,N-dimethyl-4-(methylsulfinyl)-2-[4-(trifluoromethoxy)phenoxy]- (9CI) (CA INDEX NAME)



RN 364323-05-7 HCAPLUS
 CN Benzenemethanamine, N,N-dimethyl-4-(methylsulfonyl)-2-[4-(trifluoromethoxy)phenoxy]- (9CI) (CA INDEX NAME)



REFERENCE COUNT: 7 THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L51 ANSWER 18 OF 44 HCAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: 2001:693325 HCAPLUS

DOCUMENT NUMBER: 135:257243

TITLE: Preparation of condensed imidazoles as histamine H3

INVENTOR(S): receptor ligands
 Andersen, Knud Erik; Doerwald, Florencio Zaragoza;
 Sidelmann, Ulla Grove; Rudolf, Klaus; Stenkamp, Dirk;
 Hurnaus, Rudolf; Mueller, Stephan Georg; Krist, Bernd;
 Eriksen, Birgitte; Pesche, Bernd

PATENT ASSIGNEE(S): Novo Nordisk A/S, Den.; Boehringer Ingelheim
 International G.m.b.H.

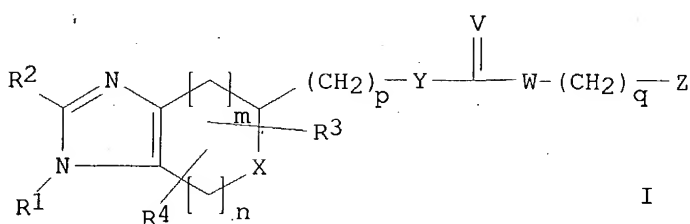
SOURCE: PCT Int. Appl., 170 pp.
 CODEN: PIXXD2

DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2001068652	A1	20010920	WO 2001-DK188	20010316
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM				
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG				
US 2002058659	A1	20020516	US 2001-810237	20010316
US 6437147	B2	20020820		
EP 1268484	A1	20030102	EP 2001-916934	20010316
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR				
JP 2003527395	T2	20030916	JP 2001-567743	20010316
US 2003135056	A1	20030717	US 2002-201865	20020723
PRIORITY APPLN. INFO.:			DK 2000-441	A 20000317
			DK 2000-1016	A 20000629
			US 2000-193741P	P 20000331
			US 2000-216553P	P 20000707
			US 2001-810237	A1 20010316
			WO 2001-DK188	W 20010316

OTHER SOURCE(S): MARPAT 135:257243
 GI



AB A novel class of imidazo heterocyclic compds. (shown as I (e.g.
 4,5,6,7-tetrahydro-1H-benzimidazole-5-carboxylic acid [(1S)-(naphth-1-

yl)ethyl]amide) as well as any optical or geometric isomer or tautomeric form thereof including mixts. of these or a pharmaceutically acceptable salt thereof), pharmaceutical compns. comprising them and use thereof in the treatment and/or prevention of diseases and disorders related to the histamine H3 receptor. In I: R1 is H or a functional group, which can be converted to H in vivo. R2 is H, C1-6-alkyl, C1-6-alkoxy, C1-6-alkylthio, halogen, cyano, trifluoromethyl, hydroxy, thiol or amino. R3 and R4 independently are H or C1-6-alkyl, which is optionally substituted with aryl or heteroaryl, which are optionally substituted with one or more substituents selected from nitro, -NR7R8, -S(O)2NR7R8, -C(O)NR7R8, hydroxy, halogen, cyano, trifluoromethyl, -OCF3, -OCHF2, -OCH2CHF2, C1-6-alkyl, C2-6-alkenyl, C2-6-alkynyl, C1-6-alkoxy, C1-6-alkylthio, C1-6-alkylsulfonyl, -C(O)OR7, C1-6-alkylcarbonyl, -C(:NOR7)C1-6-alkyl, C3-10-cycloalkyl, C3-10-cycloalkylcarbonyl, -C(:NOR7)C3-10-cycloalkyl, aryl-C1-6-alkyl, heteroaryl-C1-6-alkyl, arylamino, heteroarylarnino, aroyl, heteroaroyl, arylsulfonyl, heteroarylsulfonyl, -C(:NOR7)aryl, -C(:NOR7)heteroaryl, arylthio, heteroarylthio, aryloxy and heteroaryloxy. R7 and R8 independently are H or C1-6-alkyl. M is 0-2; n is 1-4; X is a valence bond, -O-, -S-, -S(O)-, -S(O)2- or -CF2-; p is 0-3; Y is valence bond, -O-, -S-, or -NR9-, wherein R9 is H or C1-6-alkyl; V is :O, :S, :NR10 (R10 = H, cyano, nitro, C1-6-alkyl); W is valence bond, -O-, -S-, -NR11- (R11 = H, C1-6-alkyl); q is 0-3. Z is heteroaryl, aryl, aryloxy, C3-10-cycloalkyl, C3-8-heterocyclyl or aryl annulated with C3-8-heterocyclyl, C1-6-alkyl, C2-6-alkenyl or C2-6-alkynyl, which are optionally substituted with various provisos. More particularly, the compds. are useful for the treatment and/or prevention of diseases and disorders in which an interaction with the histamine H3 receptor is beneficial. The claimed compds. generally show a high binding affinity to the histamine H3 receptor, most preferably IC50 < 500 nM. Ninety-two example preps. are included, but the methods of preparation are not claimed. Pharmaceutical compns. containing the compds. are claimed effective for reduction

of weight, suppression of appetite and treatment and/or prevention of eating disorders (e.g. bulimia, binge eating), impaired glucose tolerance (IGT), Type 2 diabetes, allergic rhinitis, ulcer, anorexia, diseases and disorders related to the serotonin-3 receptor (5-HT3; e.g. emesis), diseases and disorders related to the vanilloid receptor (e.g. pain, neurogenic inflammation, obesity), and diseases and disorders related to the alpha-2 adrenergic receptor (e.g. sleep inducing agent).

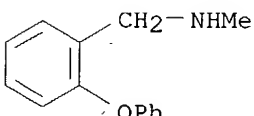
IT 361394-74-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(intermediate; preparation of condensed imidazoles as histamine H3 receptor ligands)

RN 361394-74-3 HCPLUS

CN Benzenemethanamine, N-methyl-2-phenoxy- (9CI) (CA INDEX NAME)



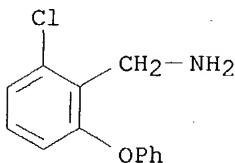
IT 175136-89-7 361394-40-3

RL: RCT (Reactant); RACT (Reactant or reagent)

(reactant; preparation of condensed imidazoles as histamine H3 receptor ligands)

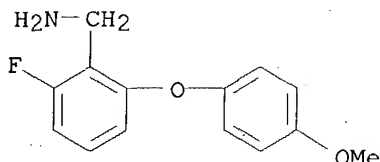
RN 175136-89-7 HCPLUS

CN Benzenemethanamine, 2-chloro-6-phenoxy- (9CI) (CA INDEX NAME)



RN 361394-40-3 HCPLUS

CN Benzenemethanamine, 2-fluoro-6-(4-methoxyphenoxy)- (9CI) (CA INDEX NAME)



REFERENCE COUNT:

6

THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L51 ANSWER 19 OF 44 HCPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: 2001:396851 HCPLUS

DOCUMENT NUMBER: 135:19652

TITLE: 3,4-Dihydro-(1H)-quinazolin-2-ones and their use as CSBP/p38 kinase inhibitors

INVENTOR(S): Adams, Jerry L.; Bower, Michael J.; Hall, Ralph F.; Griswold, Don E.; Underwood, David C.

PATENT ASSIGNEE(S): Smithkline Beecham Corporation, USA

SOURCE: PCT Int. Appl., 54 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

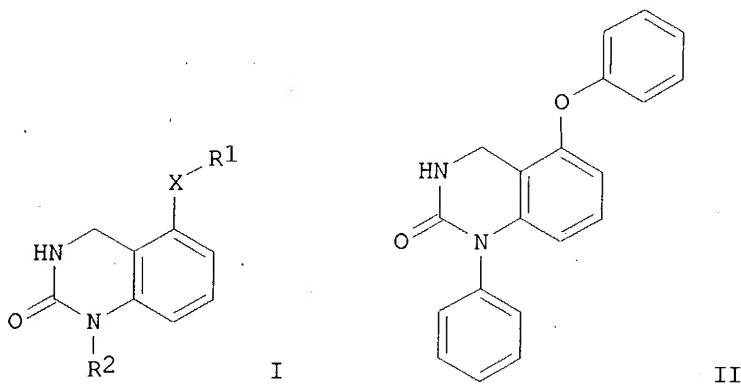
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2001038312	A1	20010531	WO 2000-US31861	20001121
W: AE, AL, AU, BA, BB, BG, BR, BZ, CA, CN, CZ, DZ, EE, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KP, KR, LC, LK, LR, LT, LV, MA, MG, MK, MN, MX, MZ, NO, NZ, PL, RO, SG, SI, SK, SL, TR, TT, TZ, UA, US, UZ, VN, YU, ZA, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM				
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG				
EP 1235814	A1	20020904	EP 2000-980569	20001121

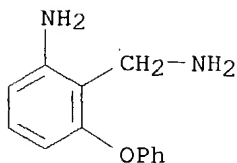
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
 IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
 JP 2003514899 T2 20030422 JP 2001-540075 20001121
 PRIORITY APPLN. INFO.: US 1999-166977P P 19991123
 WO 2000-US31861 W 20001121
 OTHER SOURCE(S): MARPAT 135:19652
 GI



AB Novel substituted quinazoline compds. are disclosed, specifically I [R1 = (un)substituted Ph, naphthyl, heterocyclyl or heteroaryl; R2 = (un)substituted alkyl, (hetero)aryl(alkyl), or heterocyclyl(alkyl); X = O or S] and their pharmaceutically acceptable salts. Also disclosed are pharmaceutical compns. containing I, and use of I in therapy as CSBP/RK/p38 kinase inhibitors. Applications of I as such to a wide variety of arthritic, inflammatory, proliferative, and viral conditions are specifically claimed. Also claimed is use of I in conjunction with various other drugs or drug classes. Three examples of I were prepared and specifically claimed. For instance, 2,6-difluorobenzonitrile was coupled first with aniline using NaH in DMSO, and then with phenol using NaH in THF. The resulting 2-phenoxy-6-(phenylamino)benzonitrile underwent reduction of the nitrile to aminomethyl using LiAlH₄, and the product underwent cyclocondensation with 1,1'-carbonyldimidazole, to give title compound II. Representative compds. I had IC₅₀ values < 50 μM in a CSBP/p38 kinase assay.

IT 342431-66-7P
 RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
 (intermediate; preparation of dihydroquinazolinones as CSBP/RK/p38 kinase inhibitors)

RN 342431-66-7 HCAPLUS
 CN Benzenemethanamine, 2-amino-6-phenoxy- (9CI) (CA INDEX NAME)



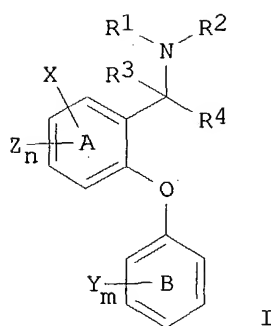
REFERENCE COUNT: 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L51 ANSWER 20 OF 44 HCAPLUS COPYRIGHT 2004 ACS on STN
 ACCESSION NUMBER: 2001:283913 HCAPLUS
 DOCUMENT NUMBER: 134:310974
 TITLE: Preparation of biaryl ether derivatives as monoamine reuptake inhibitors
 INVENTOR(S): Howard, Harry Ralph, Jr.; Adam, Mavis Diane
 PATENT ASSIGNEE(S): Pfizer Products Inc., USA
 SOURCE: PCT Int. Appl., 52 pp.
 CODEN: PIXXD2
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2001027068	A1	20010419	WO 2000-IB1373	20000927
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM				
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG				
BR 2000014733	A	20020611	BR 2000-14733	20000927
EP 1220831	A1	20020710	EP 2000-960916	20000927
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL				
JP 2003511434	T2	20030325	JP 2001-530089	20000927
EE 200200191	A	20030616	EE 2002-191	20000927
US 6410736	B1	20020625	US 2000-692335	20001019
NO 2002001659	A	20020408	NO 2002-1659	20020408
BG 106603	A	20021229	BG 2002-106603	20020410
ZA 2002002804	A	20030410	ZA 2002-2804	20020410
HR 2002000324	A1	20030831	HR 2002-324	20020412
US 2003055038	A1	20030320	US 2002-153308	20020522
US 6596741	B2	20030722		
PRIORITY APPLN. INFO.:			US 1999-159276P	P 19991013
			US 1999-167761P	P 19991129
			WO 2000-IB1373	W 20000927
			US 2000-692335	A3 20001019

OTHER SOURCE(S): MARPAT 134:310974

GI



AB The title compds. [I; rings A and B can be replaced by naphthyl group; n, m = 1-3; R1, R2 = H, alkyl, alkenyl, etc.; NR1R2 = 4-8 membered saturated (un)substituted ring containing 1-2 heteroatoms, including N atom to which R1 and R2 are attached; R3, R4 = H, alkyl optionally substituted with 1-3 F atoms; CR3R4 = 4-8 membered saturated (un)substituted carbocyclic ring; NR2CR3 = 4-8 membered saturated (un)substituted ring containing 1-2 heteroatoms, including N atom to which R2 is attached; X = (un)substituted Ph, heteroaryl, heterocyclyl; Y = H, halo, alkyl optionally substituted with 1-3 F atoms, etc.; Z = H, halo, alkoxy, etc.] and their pharmaceutically acceptable salts which exhibit activity as serotonin, norepinephrine, and dopamine reuptake inhibitors and can be used in the treatment of central nervous system and other disorders, were prepared E.g., a 3-step synthesis of I [R1 = Me; R2-R4 = H; X = 5-Ph; Z = H; Y = 3,4-Cl₂] was given. All exemplified compds. I showed IC₅₀ of ≤ 250 nM for serotonin reuptake inhibition, and IC₅₀ of ≤ 1000 nM for dopamine and for norepinephrine reuptake inhibition.

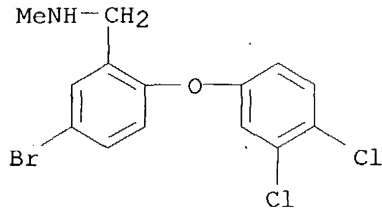
IT **289717-61-9P**

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses)

(preparation of biaryl ether derivs. as monoamine reuptake inhibitors)

RN 289717-61-9 HCAPLUS

CN Benzenemethanamine, 5-bromo-2-(3,4-dichlorophenoxy)-N-methyl- (9CI) (CA INDEX NAME)

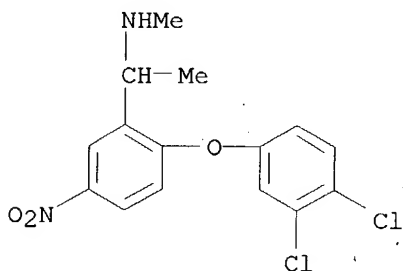


IT 334980-91-5P 334980-95-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
 (preparation of biaryl ether derivs. as monoamine reuptake inhibitors)

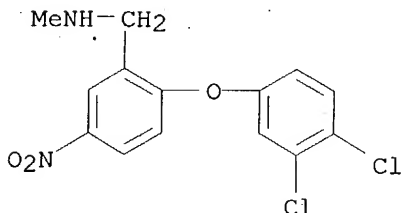
RN 334980-91-5 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N, α -dimethyl-5-nitro-
 (9CI) (CA INDEX NAME)



RN 334980-95-9 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-5-nitro- (9CI) (CA INDEX NAME)



REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L51 ANSWER 21 OF 44 HCPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: 2000:790459 HCPLUS

DOCUMENT NUMBER: 133:350040

TITLE: Preparation of agents for serotonin transporter SPECT imaging

INVENTOR(S): Kung, Hank F.

PATENT ASSIGNEE(S): USA

SOURCE: PCT Int. Appl., 76 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

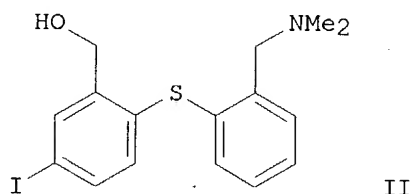
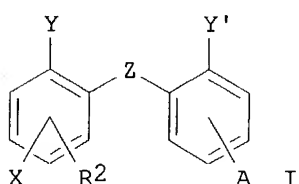
PATENT INFORMATION:

PATENT NO.

KIND DATE

APPLICATION NO. DATE

WO 2000066537	A1	20001109	WO 1999-US9344	19990430
W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM				
RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG				
AU 9936712	A1	20001117	AU 1999-36712	19990430
EP 1175388	A1	20020130	EP 1999-918906	19990430
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO				
JP 2002543173	T2	20021217	JP 2000-615370	19990430
PRIORITY APPLN. INFO.:			WO 1999-US9344	A 19990430
OTHER SOURCE(S):		MARPAT 133:350040		
GI				



AB Title compds. [I; A = H, Cl, Br, iodo, LR; L = bond, $(CH_2)_n$, $(CH_2)_nCO$; R = tetradentate ligand capable of chelating a metal(sic); R2 = H or Me; X = Cl, Br, iodo, NO₂, NR₃R₄, LR; R₃, R₄ = H, OH, alkyl, alkanoyl, etc.; 1 of Y, Y' = CH₂NR₁Me and the other = H, NO₂, CH₂OR₅, CH₂NR₁Me, NR₃R₄, LR; Z = S, O, NR₆, CR₇R₈, CO, C(:CR₇R₈); R₁, R₅, R₆ = H, LR, alkyl, alkanoyl, Ph, naphthyl, etc.; R₇, R₈ = H, Cl, alkyl; n = 1-5], with the proviso that 1 and only 1 of A, X, Y, R₁, R₃-R₆ is LR or with the proviso that 1 or both of X or A is iodo, were prepared. Data for biol. activity of I were given. E.g., 2,5-dibromobenzoic acid was coupled with freshly prepared 2-HSC₆H₄CONMe₂ with sodium methoxide as the base to give the bromo-substituted diaryl sulfide; coupling with Bu₃SnSnBu₃ in the presence of a palladium catalyst followed by substitution of the tin with iodine to give IDAM II.

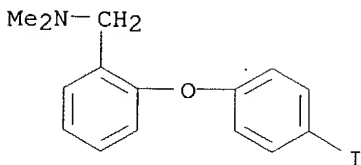
IT 305384-74-1

RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)

(preparation of agents for serotonin transporter SPECT imaging)

RN 305384-74-1 HCAPLUS

CN Benzenemethanamine, 2-(4-iodophenoxy)-N,N-dimethyl- (9CI) (CA INDEX NAME)

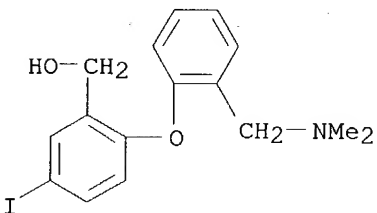


IT 291781-27-6P

RL: BPR (Biological process); BSU (Biological study, unclassified); RCT (Reactant); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); PROC (Process); RACT (Reactant or reagent)
 (preparation of agents for serotonin transporter SPECT imaging)

RN 291781-27-6 HCPLUS

CN Benzenemethanol, 2-[2-[(dimethylamino)methyl]phenoxy]-5-iodo- (9CI) (CA INDEX NAME)

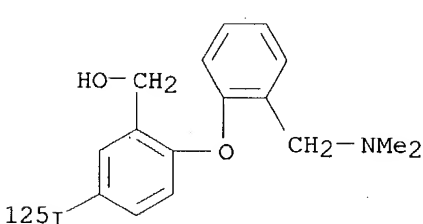


IT 291781-29-8P

RL: BPR (Biological process); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); PROC (Process)
 (preparation of agents for serotonin transporter SPECT imaging)

RN 291781-29-8 HCPLUS

CN Benzenemethanol, 2-[2-[(dimethylamino)methyl]phenoxy]-5-(iodo-125I)- (9CI) (CA INDEX NAME)



IT 291781-24-3P 291781-25-4P 291781-26-5P

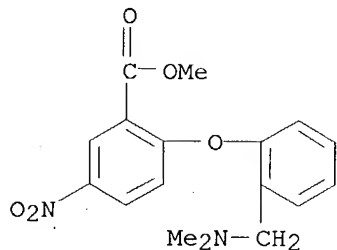
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of agents for serotonin transporter SPECT imaging)

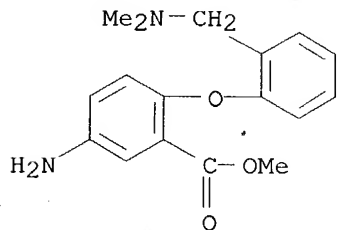
RN 291781-24-3 HCPLUS

CN Benzoic acid, 2-[2-[(dimethylamino)methyl]phenoxy]-5-nitro-, methyl ester

(9CI) (CA INDEX NAME)

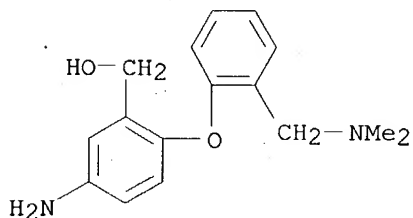


RN 291781-25-4 HCAPLUS

CN Benzoic acid, 5-amino-2-[2-[(dimethylamino)methyl]phenoxy]-, methyl ester
(9CI) (CA INDEX NAME)

RN 291781-26-5 HCAPLUS

CN Benzenemethanol, 5-amino-2-[2-[(dimethylamino)methyl]phenoxy]- (9CI) (CA INDEX NAME)



REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L51 ANSWER 22 OF 44 HCAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: 2000:761658 HCAPLUS

DOCUMENT NUMBER: 134:162754

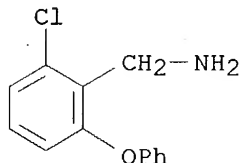
TITLE: Parallel solution-phase syntheses of functionalized bicyclo[2.2.2]octanes: generation of a library using orchestrated multi-step sequences of polymer-supported reagents and sequesterants

AUTHOR(S): Ley, Steven V.; Massi, Alessandro
 CORPORATE SOURCE: Department of Chemistry, University of Cambridge,
 Cambridge, CB2 1EW, UK
 SOURCE: Perkin 1 (2000), (21), 3645-3654
 CODEN: PERKF9; ISSN: 1470-4358
 PUBLISHER: Royal Society of Chemistry
 DOCUMENT TYPE: Journal
 LANGUAGE: English
 OTHER SOURCE(S): CASREACT 134:162754

AB An array of bicyclo[2.2.2]octane derivs. was prepared in high yield using an orchestrated multi-step sequence of polymer-supported reagents and sequestering agents, without any chromatog. purification steps. Nine intermediate libraries were synthesized, with the final library possessing five sites of diversity. Key steps included an efficient tandem Michael addition reaction of acrylates with cyclohexenones and a subsequent reductive amination reaction.

IT 175136-89-7
 RL: RCT (Reactant); RACT (Reactant or reagent)
 (parallel solution-phase syntheses of functionalized bicyclo[2.2.2]octanes)

RN 175136-89-7 HCPLUS
 CN Benzenemethanamine, 2-chloro-6-phenoxy- (9CI) (CA INDEX NAME)



REFERENCE COUNT: 58 THERE ARE 58 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L51 ANSWER 23 OF 44 HCPLUS COPYRIGHT 2004 ACS on STN
 ACCESSION NUMBER: 2000:608708 HCPLUS
 DOCUMENT NUMBER: 133:207665
 TITLE: Preparation of phenoxybenzylamines as monoamine reuptake inhibitors
 INVENTOR(S): Elliott, Mark Leonard; Howard, Harry Ralph, Jr.; Schmidt, Christopher Joseph; Seeger, Thomas Francis
 PATENT ASSIGNEE(S): Pfizer Products Inc., USA
 SOURCE: PCT Int. Appl., 60 pp.
 CODEN: PIXXD2
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 2
 PATENT INFORMATION:

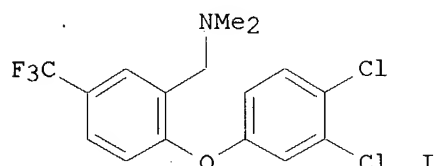
PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2000050380	A1	20000831	WO 2000-IB108	20000202
W:	AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD,			

MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK,
 SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY,
 KG, KZ, MD, RU, TJ, TM
 RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE,
 DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF,
 CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
 EP 1154984 A1 20011121 EP 2000-900785 20000202
 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
 IE, SI, LT, LV, FI, RO
 BR 2000008958 A 20011127 BR 2000-8958 20000202
 EE 200100441 A 20021216 EE 2001-441 20000202
 AU 763884 B2 20030731 AU 2000-30709 20000202
 NZ 512910 A 20031128 NZ 2000-512910 20000202
 US 2002143003 A1 20021003 US 2001-845992 20010430
 US 6677378 B2 20040113
 HR 2001000585 A1 20021231 HR 2001-585 20010807
 NO 2001004047 A 20011022 NO 2001-4047 20010820
 ZA 2001006890 A 20020923 ZA 2001-6890 20010821
 BG 105858 A 20020430 BG 2001-105858 20010830
 PRIORITY APPLN. INFO.: US 1999-121313P P 19990223
 US 2000-529207 A2 20000202
 WO 2000-IB108 W 20000202

OTHER SOURCE(S) :

MARPAT 133:207665

GI



AB ROZCR3R4NR1R2 [R = (un)substituted Ph; R1,R2 = H, alk(en)yl, alkynyl; NR1R2 = heterocyclyl; R3,R4 = H or (fluoro)alkyl; R3R4 = (un)substituted alkylene; R2R3 = atoms to complete a heterocyclic ring; Z = (un)substituted phenylene] were prepared as monoamine reuptake inhibitors (no data). Thus, 2,5-F(F₃C)C₆H₃CHO was aroxylated by 3,4-Cl₂C₆H₃OH and the product reductively aminated by Me₂NH to give title compound I.

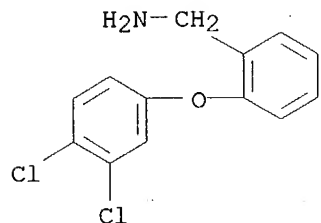
IT 146520-69-6P 146797-20-8P 289716-74-1P
 289716-75-2P 289716-79-6P 289716-80-9P
 289716-82-1P 289716-85-4P 289716-88-7P
 289716-89-8P 289716-91-2P 289716-92-3P
 289716-93-4P 289716-94-5P 289716-95-6P
 289716-96-7P 289716-97-8P 289716-98-9P
 289717-00-6P 289717-01-7P 289717-02-8P
 289717-04-0P 289717-06-2P 289717-08-4P
 289717-09-5P 289717-11-9P 289717-13-1P
 289717-15-3P 289717-16-4P 289717-17-5P
 289717-18-6P 289717-19-7P 289717-23-3P
 289717-24-4P 289717-25-5P 289717-26-6P
 289717-28-8P 289717-29-9P 289717-30-2P

289717-32-4P 289717-33-5P 289717-34-6P
 289717-35-7P 289717-36-8P 289717-37-9P
 289717-38-0P 289717-39-1P 289717-41-5P
 289717-42-6P 289717-43-7P 289717-44-8P
 289717-45-9P 289717-46-0P 289717-47-1P
 289717-48-2P 289717-49-3P 289717-50-6P
 289717-51-7P 289717-52-8P 289717-53-9P
 289717-54-0P 289717-55-1P 289717-56-2P
 289717-57-3P 289717-58-4P 289717-59-5P
 289717-60-8P 289717-61-9P 289717-62-0P
 289717-63-1P 289717-64-2P 289717-65-3P
 289717-66-4P 289717-67-5P 289717-68-6P
 289717-69-7P 289717-70-0P 289717-71-1P
 289717-72-2P 289717-73-3P 289717-74-4P
 289717-75-5P 289719-21-7P 289719-22-8P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
 (preparation of phenoxybenzylamines as monoamine reuptake inhibitors)

RN 146520-69-6 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

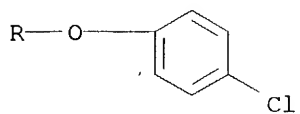
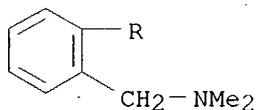
RN 146797-20-8 HCPLUS

CN Benzenemethanamine, 2-(4-chlorophenoxy)-N,N-dimethyl-, (2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 146797-19-5

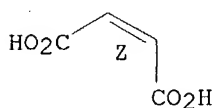
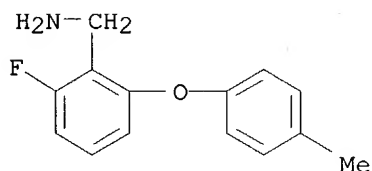
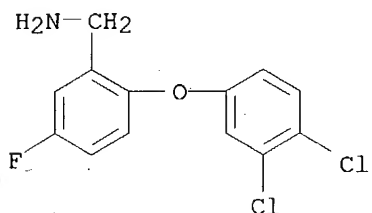
CMF C15 H16 Cl N O



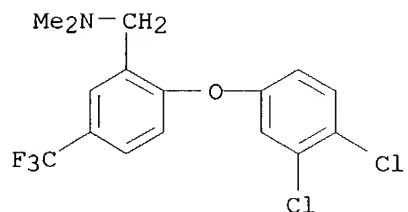
CM 2

CRN 110-16-7
CMF C₄ H₄ O₄

Double bond geometry as shown.

RN 289716-74-1 HCPLUS
CN Benzenemethanamine, 2-fluoro-6-(4-methylphenoxy)- (9CI) (CA INDEX NAME)RN 289716-75-2 HCPLUS
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro- (9CI) (CA INDEX NAME)

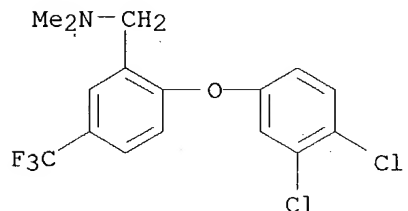
RN 289716-79-6 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-5-(trifluoromethyl)- (9CI) (CA INDEX NAME)



RN 289716-80-9 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-5-(trifluoromethyl)-, (2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

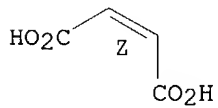
CRN 289716-79-6
 CMF C16 H14 Cl2 F3 N O



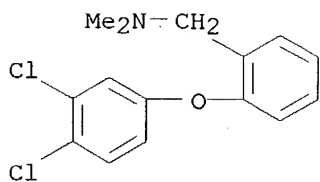
CM 2

CRN 110-16-7
 CMF C4 H4 O4

Double bond geometry as shown.



RN 289716-82-1 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-, hydrochloride (9CI) (CA INDEX NAME)

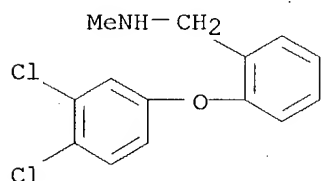


● HCl

RN 289716-85-4 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-, (2Z)-2-butenedioate
 (4:3) (9CI) (CA INDEX NAME)

CM 1

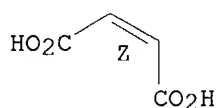
CRN 289716-84-3
 CMF C14 H13 Cl2 N O



CM 2

CRN 110-16-7
 CMF C4 H4 O4

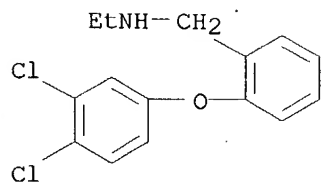
Double bond geometry as shown.



RN 289716-88-7 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-ethyl-, (2Z)-2-butenedioate
 (1:1) (9CI) (CA INDEX NAME)

CM 1

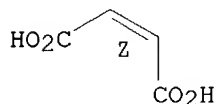
CRN 289716-87-6
 CMF C15 H15 Cl2 N O



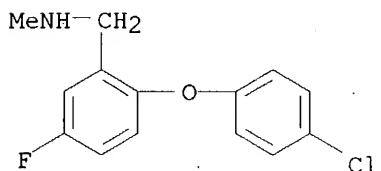
CM 2

CRN 110-16-7
CMF C₁₄ H₁₄ Cl N O₄

Double bond geometry as shown.



RN 289716-89-8 HCPLUS
 CN Benzenemethanamine, 2-(4-chlorophenoxy)-5-fluoro-N-methyl-, hydrochloride
 (9CI) (CA INDEX NAME)

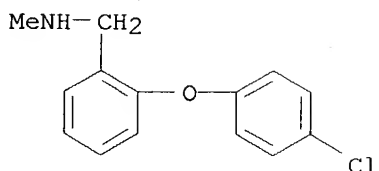


● HCl

RN 289716-91-2 HCPLUS
 CN Benzenemethanamine, 2-(4-chlorophenoxy)-N-methyl-, (2Z)-2-butenedioate
 (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 289716-90-1
CMF C₁₄ H₁₄ Cl N O

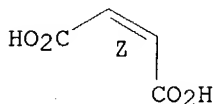


CM 2

CRN 110-16-7

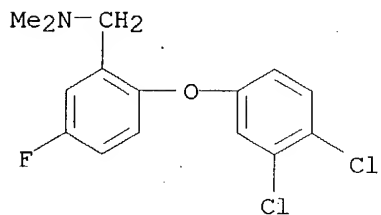
CMF C4 H4 O4

Double bond geometry as shown.



RN 289716-92-3 HCPLUS

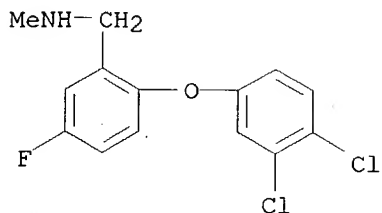
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N,N-dimethyl-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

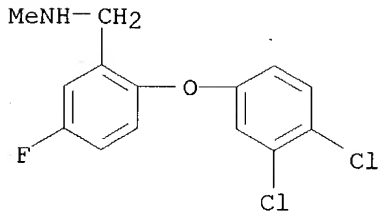
RN 289716-93-4 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N-methyl-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

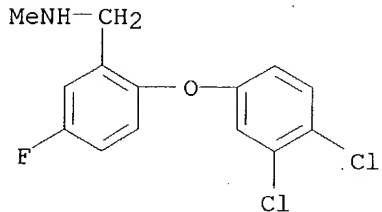
RN 289716-94-5 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N-methyl- (9CI) (CA INDEX NAME)



RN 289716-95-6 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N-methyl-, (2Z)-2-butenedioate (2:1) (9CI) (CA INDEX NAME)

CM 1

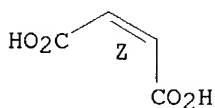
CRN 289716-94-5
 CMF C14 H12 Cl2 F N O



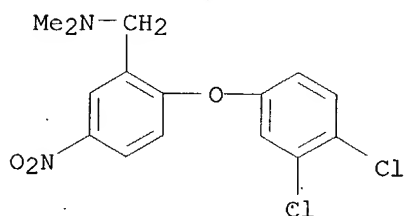
CM 2

CRN 110-16-7
 CMF C4 H4 O4

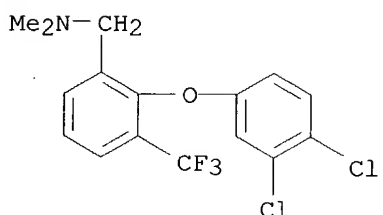
Double bond geometry as shown.



RN 289716-96-7 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-5-nitro- (9CI)
 (CA INDEX NAME)

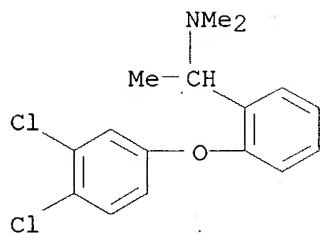


RN 289716-97-8 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-3-(trifluoromethyl)-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

RN 289716-98-9 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N, α -trimethyl-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

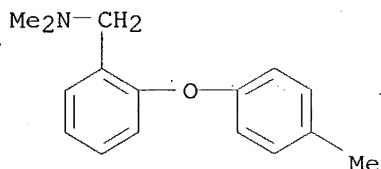
RN 289717-00-6 HCPLUS

CN Benzenemethanamine, N,N-dimethyl-2-(4-methylphenoxy)-, (2Z)-2-butenedioate
(1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 289716-99-0

CMF C16 H19 N O

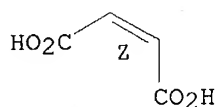


CM 2

CRN 110-16-7

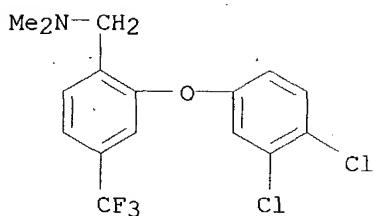
CMF C4 H4 O4

Double bond geometry as shown.



RN 289717-01-7 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-4-(trifluoromethyl)- (9CI) (CA INDEX NAME)



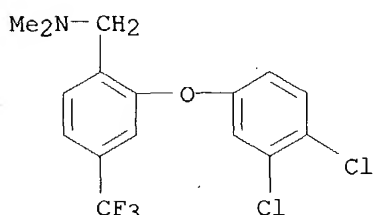
RN 289717-02-8 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-4-(trifluoromethyl)-, (2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 289717-01-7

CMF C16 H14 Cl2 F3 N O

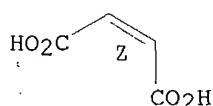


CM 2

CRN 110-16-7

CMF C4 H4 O4

Double bond geometry as shown.



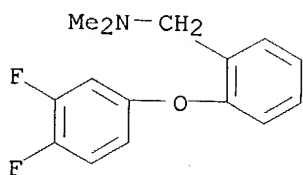
RN 289717-04-0 HCPLUS

CN Benzenemethanamine, 2-(3,4-difluorophenoxy)-N,N-dimethyl-, (2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 289717-03-9

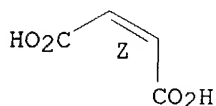
CMF C15 H15 F2 N O



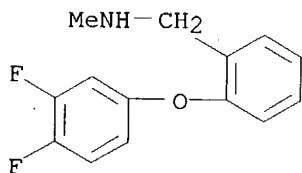
CM 2

CRN 110-16-7
CMF C4 H4 O4

Double bond geometry as shown.

RN 289717-06-2 HCPLUS
CN Benzenemethanamine, 2-(3,4-difluorophenoxy)-N-methyl-, (2Z)-2-butenedioate
(1:1) (9CI) (CA INDEX NAME)

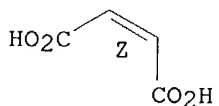
CM 1

CRN 289717-05-1
CMF C14 H13 F2 N O

CM 2

CRN 110-16-7
CMF C4 H4 O4

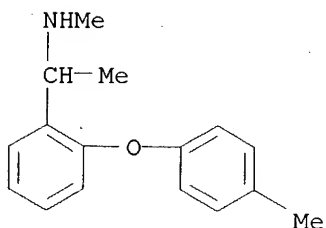
Double bond geometry as shown.



RN 289717-08-4 HCPLUS
 CN Benzenemethanamine, N, α -dimethyl-2-(4-methylphenoxy)-,
 (2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

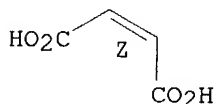
CRN 289717-07-3
 CMF C16 H19 N O



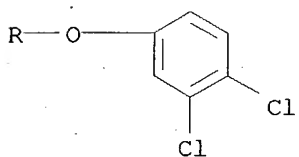
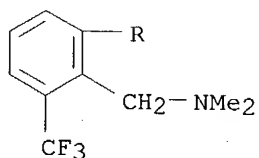
CM 2

CRN 110-16-7
 CMF C4 H4 O4

Double bond geometry as shown.



RN 289717-09-5 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N-dimethyl-6-(trifluoromethyl)-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

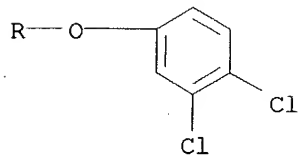
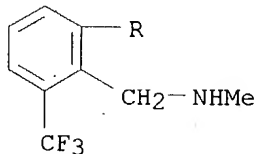
RN 289717-11-9 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-6-(trifluoromethyl)-, (2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 289717-10-8

CMF C15 H12 Cl2 F3 N O

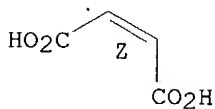


CM 2

CRN 110-16-7

CMF C4 H4 O4

Double bond geometry as shown.



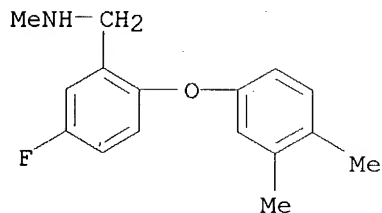
RN 289717-13-1 HCAPLUS

CN Benzenemethanamine, 2-(3,4-dimethylphenoxy)-5-fluoro-N-methyl-,
(2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 289717-12-0

CMF C16 H18 F N O

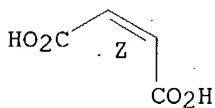


CM 2

CRN 110-16-7

CMF C4 H4 O4

Double bond geometry as shown.



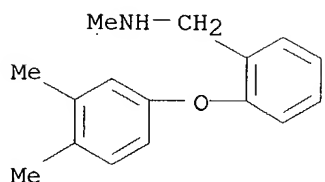
RN 289717-15-3 HCAPLUS

CN Benzenemethanamine, 2-(3,4-dimethylphenoxy)-N-methyl-, (2Z)-2-butenedioate
(1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 289717-14-2

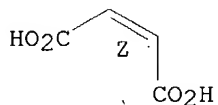
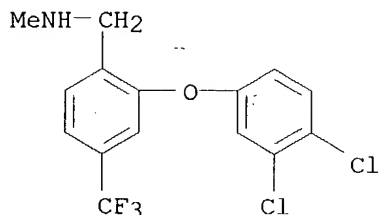
CMF C16 H19 N O



CM 2

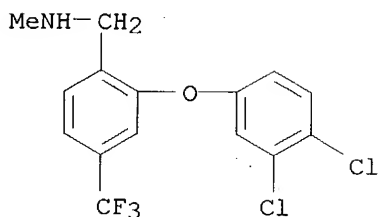
CRN 110-16-7
CMF C4 H4 O4

Double bond geometry as shown.

RN 289717-16-4 HCPLUS
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-4-(trifluoromethyl)-
(9CI) (CA INDEX NAME)RN 289717-17-5 HCPLUS
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-4-(trifluoromethyl)-
(2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

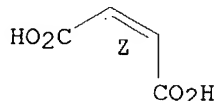
CRN 289717-16-4
CMF C15 H12 Cl2 F3 N O



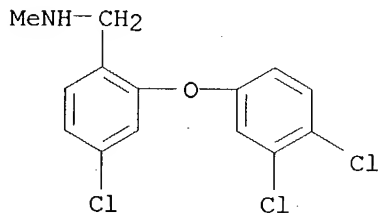
CM 2

CRN 110-16-7
CMF C₁₄ H₁₂ Cl₃ N O₄

Double bond geometry as shown.



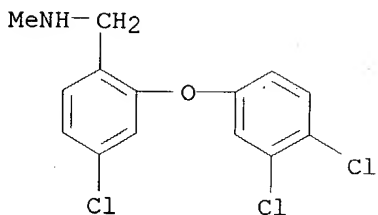
RN 289717-18-6 HCPLUS
 CN Benzenemethanamine, 4-chloro-2-(3,4-dichlorophenoxy)-N-methyl- (9CI) (CA INDEX NAME)



RN 289717-19-7 HCPLUS
 CN Benzenemethanamine, 4-chloro-2-(3,4-dichlorophenoxy)-N-methyl-, (2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

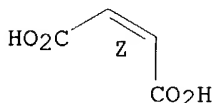
CRN 289717-18-6
CMF C₁₄ H₁₂ Cl₃ N O



CM 2

CRN 110-16-7
CMF C₄ H₄ O₄

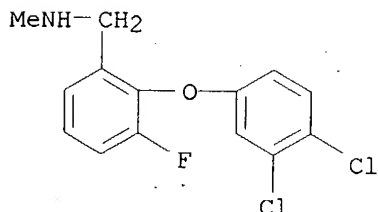
Double bond geometry as shown.



RN 289717-23-3 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-3-fluoro-N-methyl-,
(2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

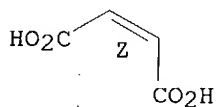
CM 1

CRN 289717-22-2
CMF C₁₄ H₁₂ Cl₂ F N O

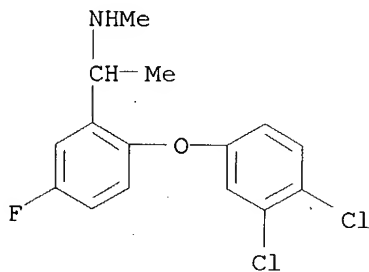
CM 2

CRN 110-16-7
CMF C₄ H₄ O₄

Double bond geometry as shown.



RN 289717-24-4 HCAPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N, α -dimethyl-
(9CI) (CA INDEX NAME)

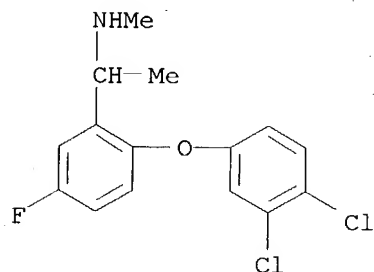
RN 289717-25-5 HCAPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N, α -dimethyl-,
(2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 289717-24-4

CMF C15 H14 Cl2 F N O

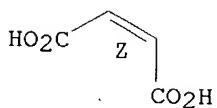


CM 2

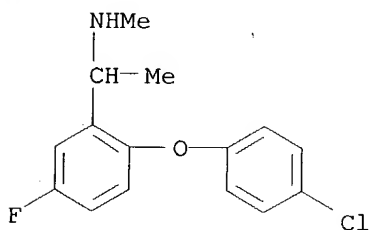
CRN 110-16-7

CMF C4 H4 O4

Double bond geometry as shown.



RN 289717-26-6 HCAPLUS
 CN Benzenemethanamine, 2-(4-chlorophenoxy)-5-fluoro-N, α -dimethyl-, hydrochloride (9CI) (CA INDEX NAME)

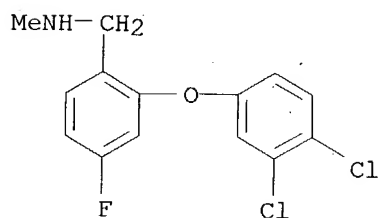


● HCl

RN 289717-28-8 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4-fluoro-N-methyl-, (2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

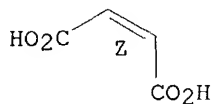
CRN 289717-27-7
 CMF C14 H12 Cl2 F N O



CM 2

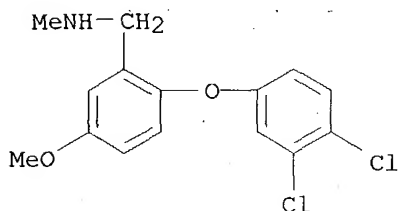
CRN 110-16-7
 CMF C4 H4 O4

Double bond geometry as shown.



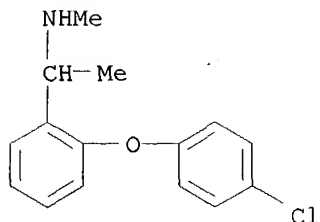
RN 289717-29-9 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-methoxy-N-methyl-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

RN 289717-30-2 HCPLUS

CN Benzenemethanamine, 2-(4-chlorophenoxy)-N, α -dimethyl-, hydrochloride (9CI) (CA INDEX NAME)

● HCl

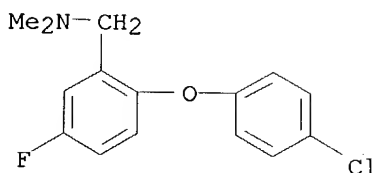
RN 289717-32-4 HCPLUS

CN Benzenemethanamine, 2-(4-chlorophenoxy)-5-fluoro-N,N-dimethyl-, (Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 289717-31-3

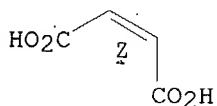
CMF C15 H15 Cl F N O



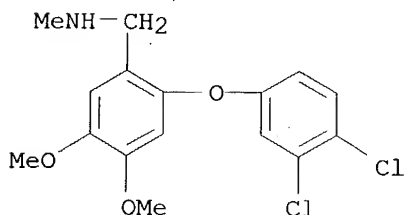
CM 2

CRN 110-16-7
CMF C₄ H₄ O₄

Double bond geometry as shown.

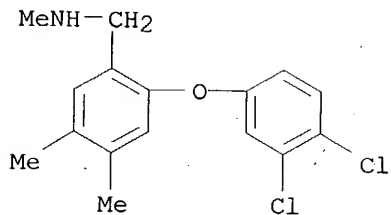


RN 289717-33-5 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4,5-dimethoxy-N-methyl-,
 hydrochloride (9CI) (CA INDEX NAME)



● HCl

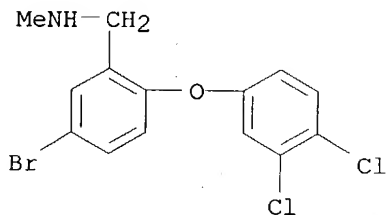
RN 289717-34-6 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,4,5-trimethyl-,
 hydrochloride (9CI) (CA INDEX NAME)



● HCl

RN 289717-35-7 HCPLUS

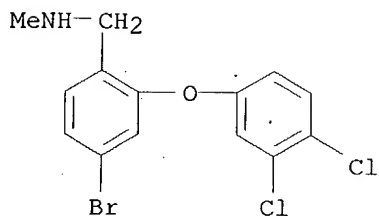
CN Benzenemethanamine, 5-bromo-2-(3,4-dichlorophenoxy)-N-methyl-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

RN 289717-36-8 HCPLUS

CN Benzenemethanamine, 4-bromo-2-(3,4-dichlorophenoxy)-N-methyl-, hydrochloride (9CI) (CA INDEX NAME)

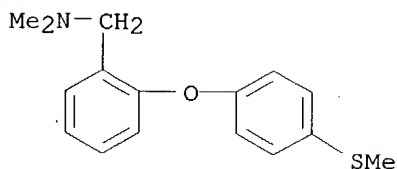


● HCl

RN 289717-37-9 HCPLUS

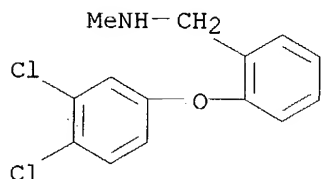
CN Benzenemethanamine, N,N-dimethyl-2-[4-(methylthio)phenoxy]-, hydrochloride

(9CI) (CA INDEX NAME)



● HCl

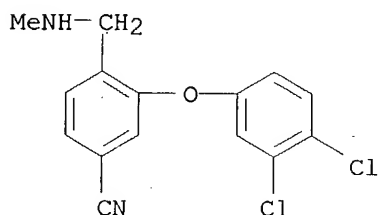
RN 289717-38-0 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-, hydrochloride (9CI)
(CA INDEX NAME)

● HCl

RN 289717-39-1 HCPLUS

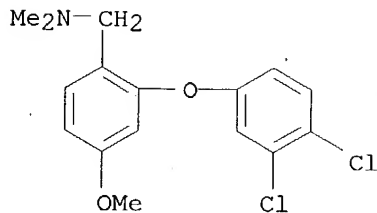
CN Benzonitrile, 3-(3,4-dichlorophenoxy)-4-[(methylamino)methyl]-, monohydrochloride (9CI) (CA INDEX NAME)



● HCl

RN 289717-41-5 HCPLUS

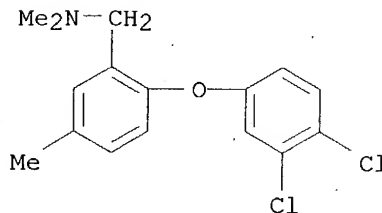
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4-methoxy-N,N-dimethyl-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

RN 289717-42-6 HCPLUS

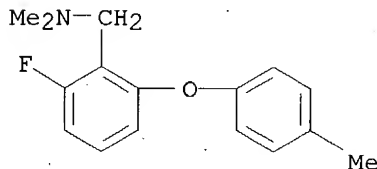
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N,5-trimethyl-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

RN 289717-43-7 HCPLUS

CN Benzenemethanamine, 2-fluoro-N,N-dimethyl-6-(4-methylphenoxy)- (9CI) (CA INDEX NAME)

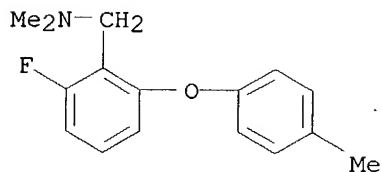


RN 289717-44-8 HCPLUS

CN Benzenemethanamine, 2-fluoro-N,N-dimethyl-6-(4-methylphenoxy)-, (2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

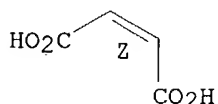
CRN 289717-43-7
 CMF C16 H18 F N O



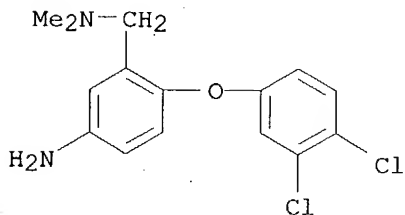
CM 2

CRN 110-16-7
 CMF C4 H4 O4

Double bond geometry as shown.

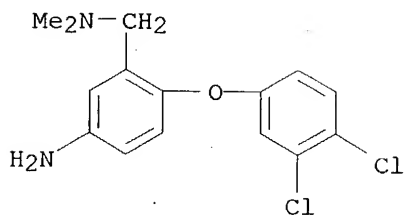


RN 289717-45-9 HCPLUS
 CN Benzenemethanamine, 5-amino-2-(3,4-dichlorophenoxy)-N,N-dimethyl-,
 monohydrochloride (9CI) (CA INDEX NAME)



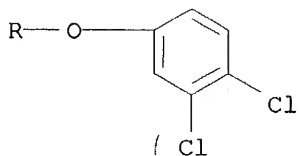
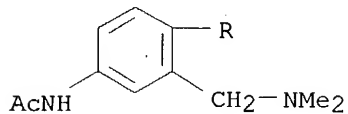
● HCl

RN 289717-46-0 HCPLUS
 CN Benzenemethanamine, 5-amino-2-(3,4-dichlorophenoxy)-N,N-dimethyl- (9CI)
 (CA INDEX NAME)



RN 289717-47-1 HCPLUS

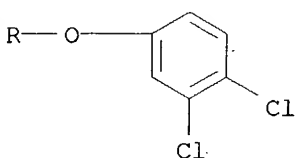
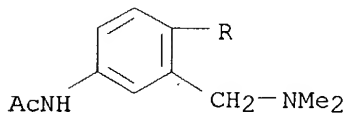
CN Acetamide, N-[4-(3,4-dichlorophenoxy)-3-[(dimethylamino)methyl]phenyl]-, monohydrochloride (9CI) (CA INDEX NAME)



● HCl

RN 289717-48-2 HCPLUS

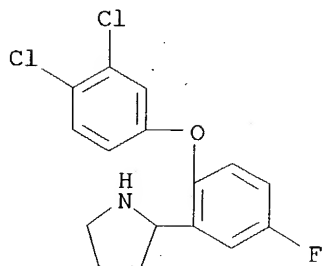
CN Acetamide, N-[4-(3,4-dichlorophenoxy)-3-[(dimethylamino)methyl]phenyl]- (9CI) (CA INDEX NAME)



RN 289717-49-3 HCPLUS

CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-, hydrochloride

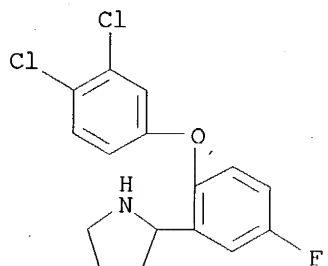
(9CI) (CA INDEX NAME)



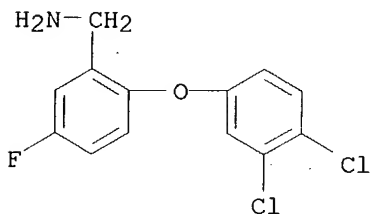
● HCl

RN 289717-50-6 HCPLUS

CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]- (9CI) (CA INDEX NAME)

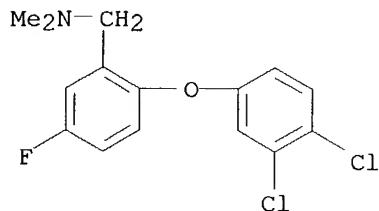


RN 289717-51-7 HCPLUS

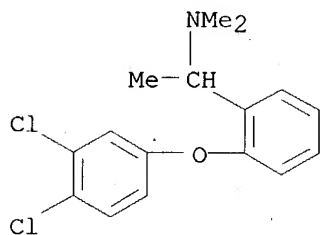
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-, hydrochloride (9CI)
(CA INDEX NAME)

● HCl

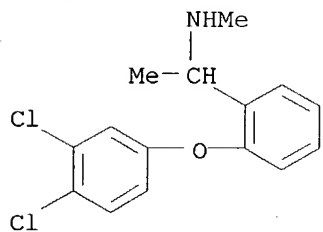
RN 289717-52-8 HCAPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N,N-dimethyl- (9CI)
(CA INDEX NAME)

RN 289717-53-9 HCAPLUS

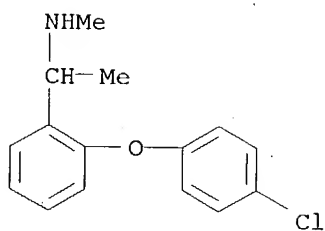
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N, α -trimethyl- (9CI)
(CA INDEX NAME)

RN 289717-54-0 HCAPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N, α -dimethyl- (9CI) (CA
INDEX NAME)

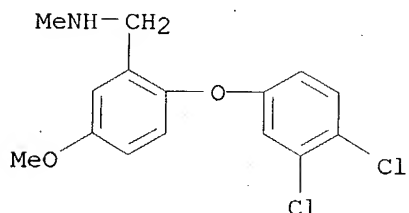
RN 289717-55-1 HCAPLUS

CN Benzenemethanamine, 2-(4-chlorophenoxy)-N, α -dimethyl- (9CI) (CA
INDEX NAME)



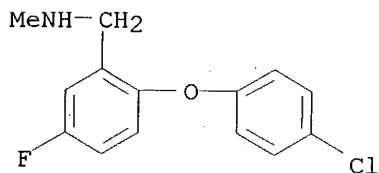
RN 289717-56-2 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-methoxy-N-methyl- (9CI) (CA INDEX NAME)



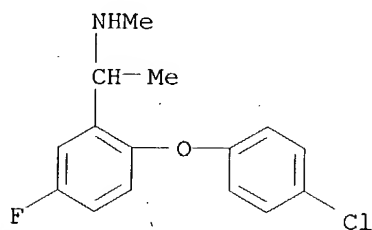
RN 289717-57-3 HCPLUS

CN Benzenemethanamine, 2-(4-chlorophenoxy)-5-fluoro-N-methyl- (9CI) (CA INDEX NAME)

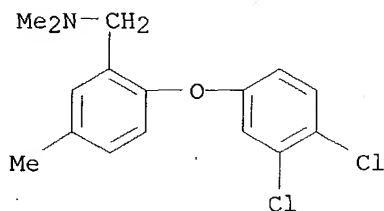


RN 289717-58-4 HCPLUS

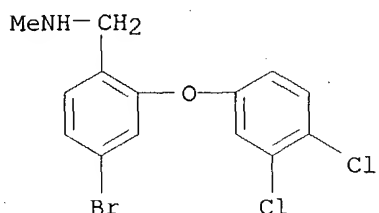
CN Benzenemethanamine, 2-(4-chlorophenoxy)-5-fluoro-N,α-dimethyl- (9CI) (CA INDEX NAME)



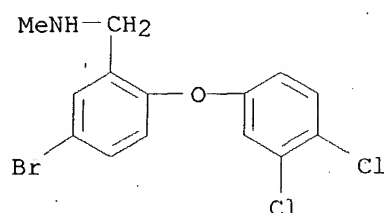
RN 289717-59-5 HCAPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,N,5-trimethyl- (9CI) (CA
INDEX NAME)

RN 289717-60-8 HCAPLUS

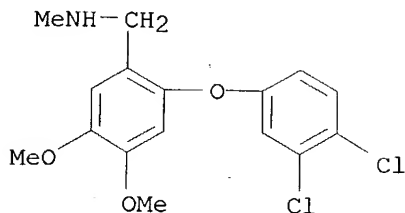
CN Benzenemethanamine, 4-bromo-2-(3,4-dichlorophenoxy)-N-methyl- (9CI) (CA
INDEX NAME)

RN 289717-61-9 HCAPLUS

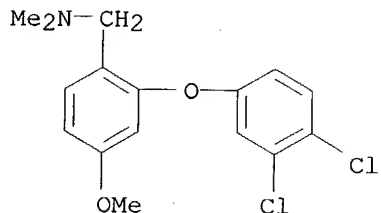
CN Benzenemethanamine, 5-bromo-2-(3,4-dichlorophenoxy)-N-methyl- (9CI) (CA
INDEX NAME)

RN 289717-62-0 HCAPLUS

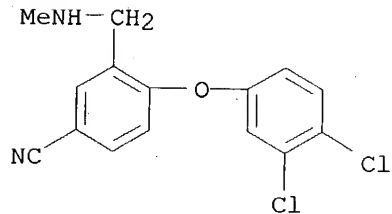
CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4,5-dimethoxy-N-methyl- (9CI)
(CA INDEX NAME)



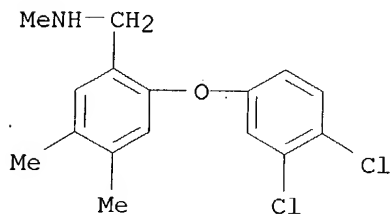
RN 289717-63-1 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4-methoxy-N,N-dimethyl- (9CI)
 (CA INDEX NAME)



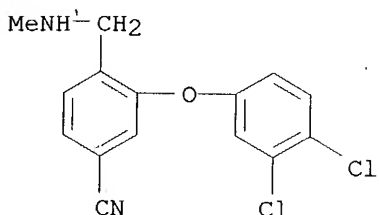
RN 289717-64-2 HCAPLUS
 CN Benzonitrile, 4-(3,4-dichlorophenoxy)-3-[(methylamino)methyl]- (9CI) (CA INDEX NAME)



RN 289717-65-3 HCAPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N,4,5-trimethyl- (9CI) (CA INDEX NAME)

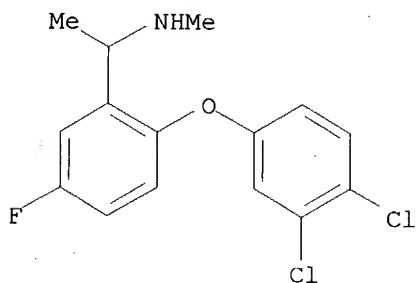


RN 289717-66-4 HCPLUS
 CN Benzonitrile, 3-(3,4-dichlorophenoxy)-4-[(methylamino)methyl]- (9CI) (CA INDEX NAME)



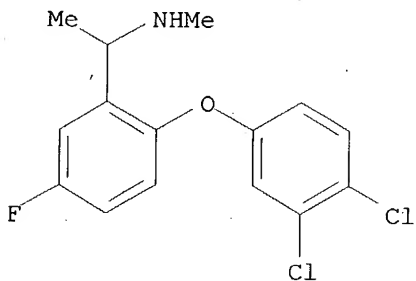
RN 289717-67-5 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N, α -dimethyl-, (+)- (9CI) (CA INDEX NAME)

Rotation (+).



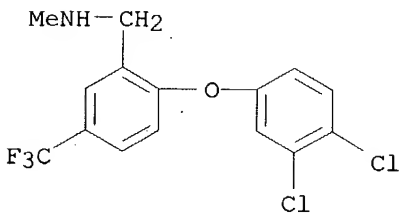
RN 289717-68-6 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-fluoro-N, α -dimethyl-, (-)- (9CI) (CA INDEX NAME)

Rotation (-).



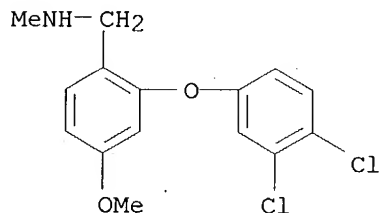
RN 289717-69-7 HCPLUS
 CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-N-methyl-5-(trifluoromethyl)-

(9CI) (CA INDEX NAME)

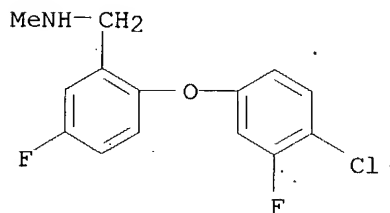


RN 289717-70-0 HCAPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4-methoxy-N-methyl- (9CI) (CA INDEX NAME)

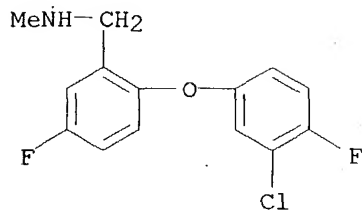


RN 289717-71-1 HCAPLUS

CN Benzenemethanamine, 2-(4-chloro-3-fluorophenoxy)-5-fluoro-N-methyl- (9CI)
(CA INDEX NAME)

RN 289717-72-2 HCAPLUS

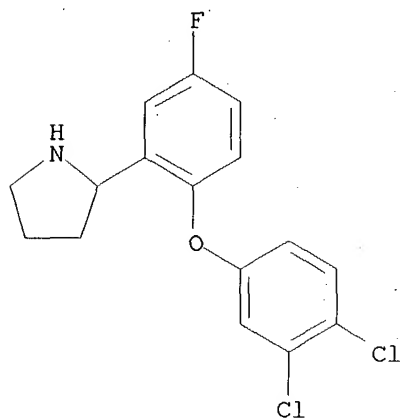
CN Benzenemethanamine, 2-(3-chloro-4-fluorophenoxy)-5-fluoro-N-methyl- (9CI)
(CA INDEX NAME)



RN 289717-73-3 HCAPLUS

CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-, (-)- (9CI) (CA INDEX NAME)

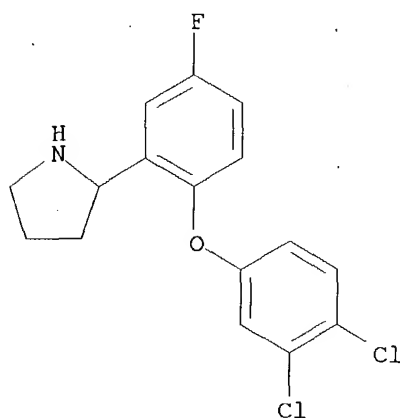
Rotation (-).



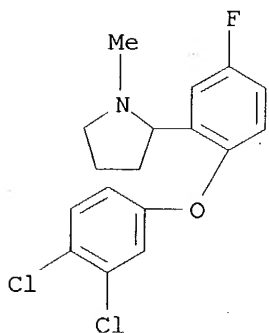
RN 289717-74-4 HCAPLUS

CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-, (+)- (9CI) (CA INDEX NAME)

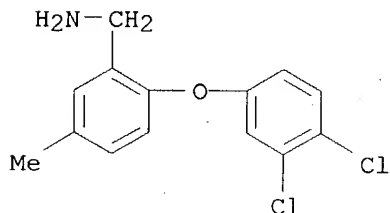
Rotation (+).



RN 289717-75-5 HCPLUS

CN Pyrrolidine, 2-[2-(3,4-dichlorophenoxy)-5-fluorophenyl]-1-methyl- (9CI)
(CA INDEX NAME)

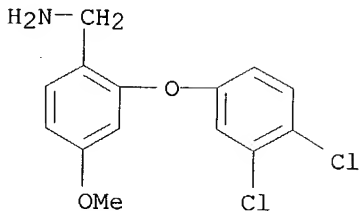
RN 289719-21-7 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-5-methyl-, hydrochloride (9CI)
(CA INDEX NAME)

● HCl

RN 289719-22-8 HCPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-4-methoxy-, hydrochloride
(9CI) (CA INDEX NAME)



● HCl

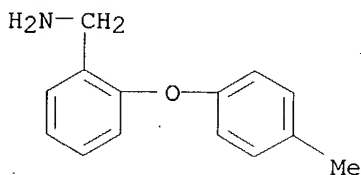
IT 289718-11-2P

RL: BYP (Byproduct); PREP (Preparation)

(preparation of phenoxybenzylamines as monoamine reuptake inhibitors)

RN 289718-11-2 HCPLUS

CN Benzenemethanamine, 2-(4-methylphenoxy)- (9CI) (CA INDEX NAME)



REFERENCE COUNT:

3

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L51 ANSWER 24 OF 44 HCPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: 2000:255979 HCPLUS

DOCUMENT NUMBER: 133:219564

TITLE: A novel serotonin transporter ligand:
(5-Iodo-2-(2-dimethylaminomethylphenoxy)-benzyl alcohol

AUTHOR(S): Zhuang, Z.-P.; Choi, S.-R.; Hou, C.; Mu, M.; Kung, M.-P.; Acton, P. D.; Kung, H. F.

CORPORATE SOURCE: Departments of Radiology and Pharmacology, University of Pennsylvania, Philadelphia, PA, USA

SOURCE: Nuclear Medicine and Biology (2000), 27(2), 169-175

CODEN: NMBIEO; ISSN: 0969-8051

PUBLISHER: Elsevier Science Inc.

DOCUMENT TYPE: Journal

LANGUAGE: English

AB The serotonin transporters (SERT) are the primary binding sites for selective serotonin reuptake inhibitors, commonly used antidepressants such as fluoxetine, sertraline, and paroxetine. Imaging of SERT with positron emission tomog. and single photon emission computed tomog. in humans would provide a useful tool for understanding how alterations of this system are related to depressive illnesses and other psychiatric disorders. In this article the synthesis and characterization of [125I]ODAM [(5-iodo-2-(2-dimethylaminomethylphenoxy)-benzyl alc.)] as an

imaging agent in the evaluation of central nervous system SERT are reported. In an initial binding study using cortical membrane homogenates of rat brain, ODAM displayed a good binding affinity with a value of $K_i = 2.8 \pm 0.88$ nM. Using LLC-PK1 cells specifically expressing the individual transporter (i.e. dopamine [DAT], norepinephrine [NET], and SERT, resp.), ODAM showed a strong inhibition on SERT ($K_i = 0.12 \pm 0.02$ nM). Inhibition consts. for the other two transporters were lower ($K_i = 3.9 \pm 0.7$ μ M and 20.0 ± 1.9 nM for DAT and NET, resp.). Initial biodistribution study in rats after an i.v. (IV) injection of [125 I]ODAM showed a rapid brain uptake and washout (2.03, 1.49, 0.79, 0.27, and 0.07% dose/organ at 2, 30, 60, 120, and 240 min, resp.). The hypothalamus region where the serotonin neurons are located exhibited a high specific uptake. Ratios of hypothalamus-cerebellum/cerebellum based on percent dose per g of these two regions showed values of 0.35, 0.86, 0.86, 0.63, and 0.34 at 2, 30, 60, 120, and 240 min, post-IV injection, resp. The specific uptake in hypothalamus can be effectively blocked by pretreatment of known SERT ligands. The results suggest that this novel ligand displays desirable in vitro and in vivo properties as a potential SERT imaging agent.

IT

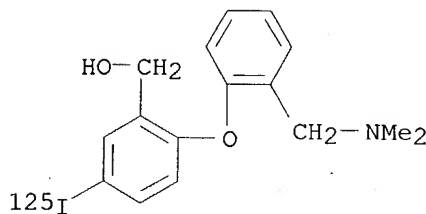
291781-29-8P

RL: BPR (Biological process); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); PROC (Process); USES (Uses)
 (preparation of radioiodinated iodo(dimethylaminomethylphenoxy)benzyl alc.
 as potential serotonin transporter imaging agent)

RN

291781-29-8 HCPLUS

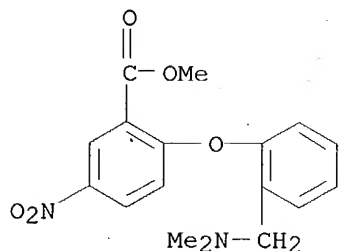
CN

Benzenemethanol, 2-[2-[(dimethylamino)methyl]phenoxy]-5-(iodo- 125 I)- (9CI)
 (CA INDEX NAME)IT **291781-24-3P 291781-25-4P 291781-26-5P****291781-27-6P**

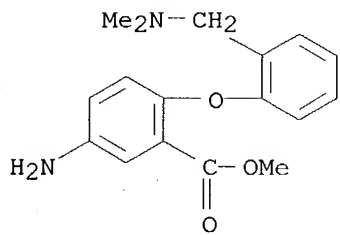
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
 (preparation of radioiodinated iodo(dimethylaminomethylphenoxy)benzyl alc.
 as potential serotonin transporter imaging agent)

RN 291781-24-3 HCPLUS

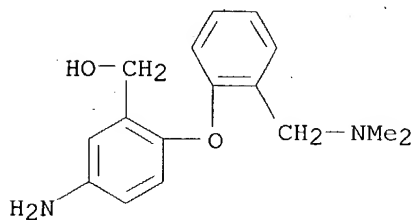
CN Benzoic acid, 2-[2-[(dimethylamino)methyl]phenoxy]-5-nitro-, methyl ester
 (9CI) (CA INDEX NAME)



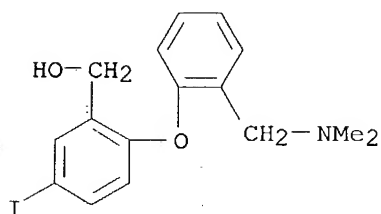
RN 291781-25-4 HCAPLUS

CN Benzoic acid, 5-amino-2-[2-[(dimethylamino)methyl]phenoxy]-, methyl ester
(9CI) (CA INDEX NAME)

RN 291781-26-5 HCAPLUS

CN Benzenemethanol, 5-amino-2-[2-[(dimethylamino)methyl]phenoxy]- (9CI) (CA
INDEX NAME)

RN 291781-27-6 HCAPLUS

CN Benzenemethanol, 2-[2-[(dimethylamino)methyl]phenoxy]-5-iodo- (9CI) (CA
INDEX NAME)

REFERENCE COUNT: 30 THERE ARE 30 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L51 ANSWER 25 OF 44 HCPLUS COPYRIGHT 2004 ACS on STN
 ACCESSION NUMBER: 1999:618468 HCPLUS
 DOCUMENT NUMBER: 132:233682
 TITLE: Single-photon emission tomography imaging of serotonin transporters in the nonhuman primate brain with [123I]ODAM
 AUTHOR(S): Acton, Paul D.; Mu, Mu; Plossl, Karl; Hou, Catherine; Siciliano, Michael; Zhuang, Zhi-Ping; Oya, Shunichi; Choi, Seok-Rye; Kung, Hank F.
 CORPORATE SOURCE: Department of Radiology, University of Pennsylvania, Philadelphia, PA, 19104, USA
 SOURCE: European Journal of Nuclear Medicine (1999), 26(10), 1359-1362
 CODEN: EJNMD9; ISSN: 0340-6997
 PUBLISHER: Springer-Verlag
 DOCUMENT TYPE: Journal
 LANGUAGE: English

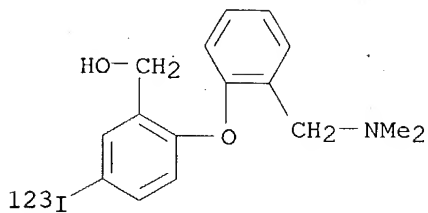
AB We have described previously a selective serotonin transporter (SERT) radioligand, [123I]IDAM. We now report a similarly potent, but more stable IDAM derivative, 5-iodo-2-[2-[(dimethylamino)methyl]phenoxy]benzyl alc. ([123I]ODAM). The imaging characteristics of this radioligand were studied and compared against [123I]IDAM. Dynamic sequences of single-photon emission tomog. (SPET) scans were obtained on three female baboons after injection of 375 MBq of [123I]ODAM. Displacing doses (1 mg/kg) of the selective SERT ligand (+)McN5652 were administered 120 min after injection of [123I]ODAM. Total integrated brain uptake of [123I]ODAM was about 30% higher than [123I]IDAM. After 60-120 min, the regional distribution of tracer within the brain reflected the characteristic distribution of SERT. Peak specific binding in the midbrain occurred 120 min after injection, with an equilibrium midbrain to cerebellar ratio of 1.50 ± 0.08 , which was slightly lower than the value for [123I]IDAM (1.80 ± 0.13). Both the binding kinetics and the metabolism of [123I]ODAM were slower than those of [123I]IDAM. Following injection of a competing SERT ligand, (+)McN5652, the tracer exhibited washout from areas with high concns. of SERT, with a dissociation kinetic rate constant $k_{off} = 0.0085 \pm 0.0028$ min⁻¹ in the midbrain. Similar studies using nisoxetine and methylphenidate showed no displacement, consistent with its low binding affinity to norepinephrine and dopamine transporters, resp. These results suggest that [123I]ODAM is suitable for selective SPET imaging of SERT in the primate brain, with higher uptake and slower kinetics and metabolism than [123I]IDAM, but also a slightly lower selectivity for SERT.

IT 262273-73-4

RL: BPR (Biological process); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses)
 (SPET imaging of serotonin transporters in nonhuman primate brain with [123I]ODAM)

RN 262273-73-4 HCPLUS

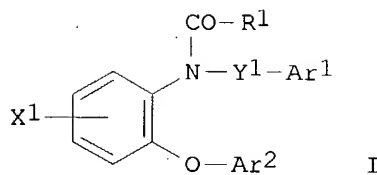
CN Benzenemethanol, 2-[2-[(dimethylamino)methyl]phenoxy]-5-(iodo-123I)- (9CI)
 (CA INDEX NAME)



REFERENCE COUNT: 9 THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L51 ANSWER 26 OF 44 HCPLUS COPYRIGHT 2004 ACS on STN
 ACCESSION NUMBER: 1999:113636 HCPLUS
 DOCUMENT NUMBER: 130:182255
 TITLE: Preparation of aryloxyaniline derivatives as therapeutic agents with high affinity for the MDR receptors
 INVENTOR(S): Nakazato, Atsuro; Okubo, Taketoshi; Nakamura, Toshio; Chaki, Shigeyuki; Tomisawa, Kazuyuki; Nagamine, Masashi; Yamamoto, Kenji; Harada, Koichiro; Yoshida, Masanori
 PATENT ASSIGNEE(S): Taisho Pharmaceutical Co., Ltd., Japan; Nihon, Nohyaku Co., Ltd.
 SOURCE: PCT Int. Appl., 59 pp.
 DOCUMENT TYPE: Patent
 LANGUAGE: Japanese
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9906353	A1	19990211	WO 1998-JP3442	19980803
W: AU, CA, CN, KR, US				
RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE				
AU 9884625	A1	19990222	AU 1998-84625	19980803
AU 729000	B2	20010125		
JP 11171844	A2	19990629	JP 1998-218784	19980803
EP 1004573	A1	20000531	EP 1998-935326	19980803
EP 1004573	B1	20021030		
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI				
AT 226932	E	20021115	AT 1998-935326	19980803
PT 1004573	T	20030331	PT 1998-98935326	19980803
ES 2186186	T3	20030501	ES 1998-935326	19980803
US 6333358	B1	20011225	US 2000-485006	20000201
US 2002147191	A1	20021010	US 2001-922807	20010807
US 6476056	B2	20021105		
PRIORITY APPLN. INFO.:			JP 1997-209123	A 19970804
			WO 1998-JP3442	W 19980803
			US 2000-485006	A3 20000201
OTHER SOURCE(S): GI		MARPAT 130:182255		



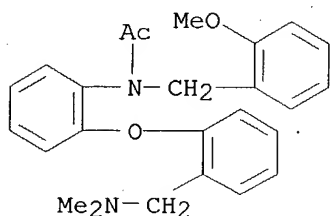
AB The title compds. I [Ar1 and Ar2 represent each a substituted or unsubstituted Ph, pyridyl, or naphthyl group; R1 represents a hydrogen atom, an alkyl group or the like; X1 represents a hydrogen atom, an alkyl group or the like; and Y1 represents a branched or unbranched 1-6 C alkylene group or a single bond] are prepared. I are useful in the treatment of anxiety and associated diseases, depression, and the like. In an in vitro test for affinity for the mitochondrial DBI receptors, the title compound I [Ar1Y1 = 3-methoxybenzyl; Ar2 = phenyl; R1 = methyl; X1 = H] showed IC50 of 1.38 nM.

IT 220553-21-9P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of aryloxyaniline derivs. as therapeutic agents with high affinity for the MDR receptors)

RN 220553-21-9 HCPLUS

CN Acetamide, N-[2-[2-[(dimethylamino)methyl]phenoxy]phenyl]-N-[(2-methoxyphenyl)methyl]- (9CI) (CA INDEX NAME)



REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L51 ANSWER 27 OF 44 HCPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: 1997:499172 HCPLUS

DOCUMENT NUMBER: 127:176352

TITLE: Quinolin-2(1H)-ones as NMDA receptor antagonists

INVENTOR(S): Ackermann, Karl-august; Gottschlich, Rudolf; Holzemann, Gunter; Leibrock, Joachim; Rautenberg, Wilfried; Seyfried, Christoph

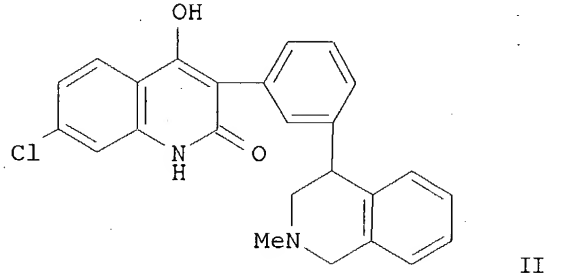
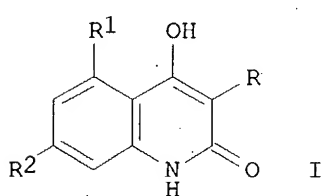
PATENT ASSIGNEE(S): Merck Patent G.m.b.H., Germany; Gottschlich, Rudolf; Holzemann, Gunter; Leibrock, Joachim; Rautenberg,

SOURCE: Wilfried; Seyfried, Christoph
 PCT Int. Appl., 43 pp.
 CODEN: PIXXD2

DOCUMENT TYPE: Patent
 LANGUAGE: German
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

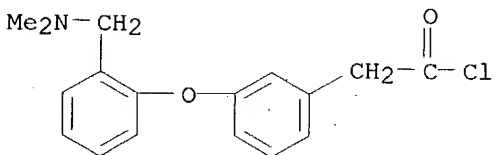
PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9726244	A1	19970724	WO 1997-EP84	19970110
W: AU, BR, CA, CN, CZ, HU, JP, KR, LT, LV, MX, NO, PL, RU, SI, SK, UA, US				
RW: AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE				
DE 19601782	A1	19970724	DE 1996-19601782	19960119
CA 2243474	AA	19970724	CA 1997-2243474	19970110
AU 9713112	A1	19970811	AU 1997-13112	19970110
AU 716230	B2	20000224		
EP 885196	A1	19981223	EP 1997-900586	19970110
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, PT, IE, SI, LT, LV, FI				
CN 1211974	A	19990324	CN 1997-192395	19970110
BR 9707027	A	19990720	BR 1997-7027	19970110
JP 2000503308	T2	20000321	JP 1997-525656	19970110
ZA 9700364	A	19970722	ZA 1997-364	19970116
NO 9803333	A	19980918	NO 1998-3333	19980717
US 6028080	A	20000222	US 1998-101837	19980717
PRIORITY APPLN. INFO.:			DE 1996-19601782 A	19960119
			WO 1997-EP84	W 19970110

OTHER SOURCE(S): MARPAT 127:176352
 GI



- AB Quinolinones I [R = substituted Ph; R1, R2 = H, halogen, alkyl, alkoxy] were prepared fo use in treating neurodegenerative disorders (no data). Thus, the quinolinone II and its enantiomers were obtained from 2-BrCH₂COC₆H₄CH₂CO₂Me in 9 steps.
- IT **193819-50-0P**
 RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
 (preparation of arylquinolinones as NMDA receptor antagonists)

RN 193819-50-0 HCAPLUS
 CN Benzeneacetyl chloride, 3-[2-[(dimethylamino)methyl]phenoxy]- (9CI) (CA
 INDEX NAME)



L51 ANSWER 28 OF 44 HCAPLUS COPYRIGHT 2004 ACS on STN
 ACCESSION NUMBER: 1997:151532 HCAPLUS
 DOCUMENT NUMBER: 126:157822
 TITLE: Synthesis of N-substituted oligomers as therapeutic agents
 INVENTOR(S): Zuckermann, Ronald N.; Goff, Dane A.; Ng, Simon;
 Spear, Kerry; Scott, Barbara O.; Sigmund, Aaron C.;
 Goldsmith, Richard A.; Marlowe, Charles K.; Pei,
 Yazhong; Richter, Lutz; Simon, Reyna
 PATENT ASSIGNEE(S): Chiron Corporation, USA
 SOURCE: PCT Int. Appl., 175 pp.
 CODEN: PIIXD2
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 3
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9640202	A1	19961219	WO 1996-US8832	19960604
W: AL, AM, AT, AU, AZ, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, EE, ES, FI, GB, GE, HU, IS, JP, KE, KG, KP, KR, KZ, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI				
RW: KE, LS, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML				
US 5877278	A	19990302	US 1995-487282	19950607
AU 9662534	A1	19961230	AU 1996-62534	19960604
EP 789577	A1	19970820	EP 1996-921278	19960604
EP 789577	B1	20030312		
R: AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LI, LU, MC, NL, PT, SE				
JP 11507049	T2	19990622	JP 1996-501317	19960604
AT 234268	E	20030315	AT 1996-921278	19960604
PRIORITY APPLN. INFO.:			US 1995-487282	A 19950607
			US 1992-950853	B2 19920924
			US 1993-126539	B2 19930924
			US 1994-277228	B2 19940718
			WO 1996-US8832	W 19960604

AB The title process comprises a solid-phase method for synthesis of N-substituted oligomers, e.g., poly(N-substituted glycines) having a wide variety of side-chain substituents, to obtain compds. of potential

therapeutic interest. Each N-substituted glycine monomer is assembled from two sub-monomers directly on the solid support. Each cycle of monomer addition consists of two steps: (1) acylation of a support-bound amine with an acylating agent containing a group capable of nucleophilic displacement by -NH₂, such as a haloacetic acid, and (2) introduction of the side-chain by nucleophilic displacement of the leaving group with a second submonomer such as a primary amine, alkoxyamine, semicarbazide, acyl hydrazide, carbazate or the like. Repetition of the two step cycle of acylation and displacement gives the desired oligomers. Combinatorial libraries are disclosed.

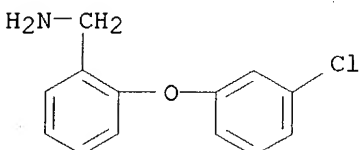
IT 186700-05-0

RL: RCT (Reactant); RACT (Reactant or reagent)

(synthesis of N-substituted oligomers as therapeutic agents)

RN 186700-05-0 HCPLUS

CN Benzenemethanamine, 2-(3-chlorophenoxy)- (9CI) (CA INDEX NAME)



L51 ANSWER 29 OF 44 HCPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: 1997:148791 HCPLUS

DOCUMENT NUMBER: 126:157515

TITLE: Preparation of ortho-substituted aromatic compounds, containing three (het)aryl moieties as prostaglandin E2-(PGE2)-antagonists

INVENTOR(S): Breault, Gloria Anne; Oldfield, John; Tucker, Howard; Warner, Peter

PATENT ASSIGNEE(S): Zeneca Limited, UK

SOURCE: Eur. Pat. Appl., 62 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent

LANGUAGE: English

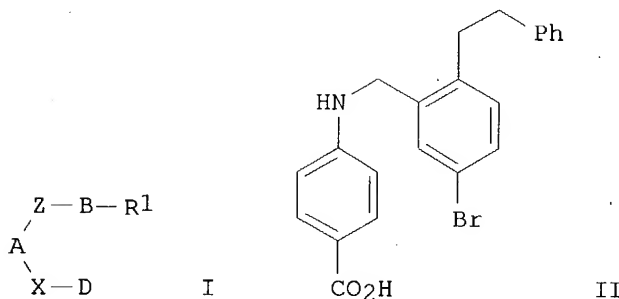
FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 752421	A1	19970108	EP 1996-304888	19960702
R: CH, DE, FR, GB, IT, LI				
US 5834468	A	19981110	US 1996-673878	19960702
JP 09040607	A2	19970210	JP 1996-173990	19960703
US 6057345	A	20000502	US 1998-183578	19981030
US 2003139418	A1	20030724	US 2002-284610	20021031
PRIORITY APPLN. INFO.:			GB 1995-13900	A 19950707
			GB 1995-13902	A 19950707
			GB 1995-13903	A 19950707
			GB 1995-13923	A 19950707
			GB 1995-13924	A 19950707
			GB 1995-13927	A 19950707
			US 1996-673878	A3 19960702

US 1998-183578 A3 19981030
 US 2000-505969 B1 20000217
 US 2001-811779 B1 20010319

OTHER SOURCE(S): MARPAT 126:157515
 GI



AB The title compds. [I; A = (un)substituted Ph, naphthyl, pyridazinyl, etc., and the -ZBR1 and -XD linking groups are positioned in a 1,2 relationship to one another on ring carbon atoms and the ring atom positioned ortho to the -X- linking group (and therefore in the 3-position relative to the -Z-linking group) is not substituted; B = (un)substituted Ph, pyridyl, thiazolyl, etc.; D = (un)substituted pyridyl, pyrazinyl, pyrimidinyl, etc.; R1 = COOH, tetrazolyl, etc., and is positioned on ring B in a 1,3 or 1,4 relationship with the -Z- linking group in 6-membered rings and in a 1,3-relationship with the -Z- linking group in 5-membered rings; X = OCH₂, SCH₂, CH₂CH₂, etc.; Z = CH(R3)CH(R3)N(R2), N(R2)CH(R3), etc. (wherein R2 = H, C1-6 alkyl, C2-6 alkenyl, etc.; R3 = H, C1-4 alkyl)] were prepared. Thus, reaction of 5-bromo-2-(phenethyl)benzaldehyde with Me 4-aminobenzoate followed by treatment of the intermediate with NaBH₄, and hydrolysis of the resulting Me 4-{N-[5-bromo-2-(phenethyl)benzyl]amino}benzoate with 2N aqueous NaOH in MeOH/THF afforded the title compound II. Compds. I are useful in the treatment of pain such as the pain associated with joint conditions (such as rheumatoid arthritis and osteoarthritis), postoperative pain, post-partum pain, the pain associated with dental conditions (such as dental caries and gingivitis), the pain associated with burns (including sunburn), the treatment of bone disorders (such as osteoporosis, hypercalcemia of malignancy and Paget's disease), the pain associated with sports injuries and sprains and all other painful conditions in which E-type prostaglandins wholly or in part play a pathophysiol. role. In general, compds. I are effective at 0.05-25 mg/kg/day.

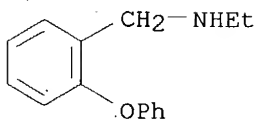
IT 186797-05-7P
 RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of ortho-substituted aromatic compds. containing three (het)aryl

moieties as prostaglandin E2-(PGE2)-antagonists)

RN 186797-05-7 HCPLUS

CN Benzenemethanamine, N-ethyl-2-phenoxy- (9CI) (CA INDEX NAME)



L51 ANSWER 30 OF 44 HCPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: 1997:44662 HCPLUS

DOCUMENT NUMBER: 126:59751

TITLE: Preparation of di- and tricarboxybenzamides and
analogs as squalene synthetase and protein
farnesyltransferase inhibitorsINVENTOR(S): Baker, William R.; Rosenberg, Saul H.; Fung, K. L.
Anthony; Rockway, Todd W.; Fakhouri, Stephen A.;
Garvey, David S.; Donner, B. Gregory; O'Connor,
Stephen J.; Prasad, Rajnandan N.; Shen, Wang; Stout,
David M.; Sullivan, Gerard M.

PATENT ASSIGNEE(S): Abbott Laboratories, USA

SOURCE: PCT Int. Appl., 241 pp.

CODEN: PIIXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

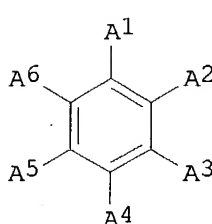
FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

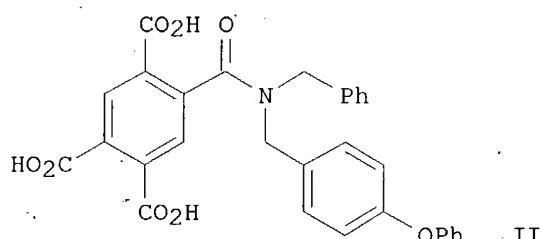
PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9634851	A1	19961107	WO 1996-US6193	19960502
W: AU, CA, JP, KR, MX				
RW: AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE				
US 5783593	A	19980721	US 1996-633262	19960429
AU 9656731	A1	19961121	AU 1996-56731	19960502
PRIORITY APPLN. INFO.:				
		US 1995-429095		19950503
		US 1996-633262		19960429
		US 1993-147708		19931104
		US 1994-289711		19940909
		US 1994-322783		19941018
		WO 1996-US6193		19960502

OTHER SOURCE(S): MARPAT 126:59751

GI



I

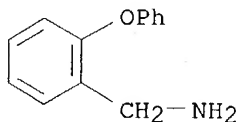


II

AB Title compds. [e.g., I; A1 = ZCONR1R2; A2,A4, and A5 or A2 and A4 or A3 and

A4 = (protected) CO₂H and the other An = H; R1 = (chloro)benzyl, (CH₂)₂-4Ph, CH₂C₆H₄(OPh)-4; R2 = (CH₂)₁₋₂C₆H₄(OPh)-4; Z = bond, NR, O; R = H, (cyclo)alkyl, aralkyl, cycloalkylalkyl] were prepared. Thus, 4-(PhO)C₆H₄CHO was reductively aminated by H₂CH₂Ph and the product amidated by 1,2,4,5-benzenetetracarboxylic dianhydride to give title compound II. Data for in vitro inhibition of protein farnesyltransferase by selected I were given.

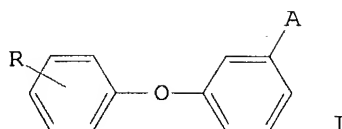
IT 107624-14-6, Benzenemethanamine, 2-phenoxy-
 RL: RCT (Reactant); RACT (Reactant or reagent)
 (preparation of di- and tricarboxybenzamides and analogs as squalene synthetase and protein farnesyltransferase inhibitors)
 RN 107624-14-6 HCPLUS
 CN Benzenemethanamine, 2-phenoxy- (9CI) (CA INDEX NAME)



L51 ANSWER 31 OF 44 HCPLUS COPYRIGHT 2004 ACS on STN
 ACCESSION NUMBER: 1993:191327 HCPLUS
 DOCUMENT NUMBER: 118:191327
 TITLE: Preparation of (aminomethylphenoxy)halobenzenes and related compounds as antidepressants
 INVENTOR(S): Ruigt, Gerardus Stephanus Franciscus; Leysen, Dirk; Wieringa, Johannes Hubertus
 PATENT ASSIGNEE(S): AKZO N. V., Neth.
 SOURCE: Eur. Pat. Appl., 8 pp.
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 516234	A1	19921202	EP 1992-201484	19920525
EP 516234	B1	19950913		
R: AT, BE, CH, DE, DK, ES, FR, ZA 9203283	A	19930127	GB, GR, IT, LI, LU, MC, NL, PT, SE ZA 1992-3283	19920506
CA 2068373	AA	19921130	CA 1992-2068373	19920511
AU 9217104	A1	19921203	AU 1992-17104	19920522
AU 650136	B2	19940609		
ES 2079783	T3	19960116	ES 1992-201484	19920525
NO 9202110	A	19921130	NO 1992-2110	19920527
NO 178395	B	19951211		
NO 178395	C	19960320		
JP 05148197	A2	19930615	JP 1992-137276	19920528
US 5190965	A	19930302	US 1992-891545	19920529
US 5430063	A	19950704	US 1993-12700	19930203
PRIORITY APPLN. INFO.:			EP 1991-201288	19910529
			US 1992-891545	19920529
OTHER SOURCE(S):	MARPAT	118:191327		

GI



AB Title compds. I ($R = 1$ or 2 halo; $A = R_2R_1NCH_2$ wherein $R_1, R_2 = H$, alkyl, $4,5$ -dihydro- $1H$ -imidazolyl), were prepared $4-ClC_6H_4OH$, NaH , 18 -crown- 6 , and $2-BrC_6H_4CN$ were heated in DMF at 100° for 16 h to give $2-(4-ClC_6H_4O)C_6H_4CN$ which was treated with $LiAlH_4$ in THF to give I ($R = 4-Cl$, $A = CH_2H_2N \cdot HCl$) (II). The $1/2$ life of II in rats was $1-2.5$ h.

IT 146797-18-4P

RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of)

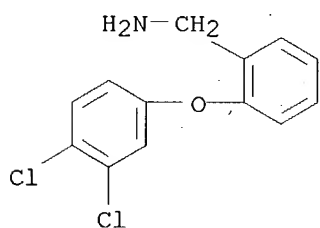
RN 146797-18-4 HCPLUS

CN 2-Naphthalenecarboxylic acid, $4,4'$ -methylenebis[3-hydroxy-, compd. with
 $2-(3,4-dichlorophenoxy)benzenemethanamine (1:2)$ (9CI) (CA INDEX NAME)

CM 1

CRN 146797-17-3

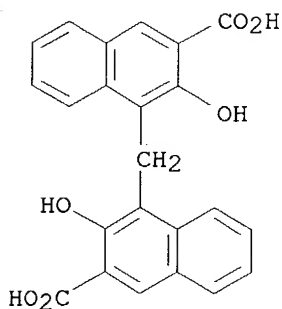
CMF C13 H11 Cl2 N O



CM 2

CRN 130-85-8

CMF C23 H16 O6

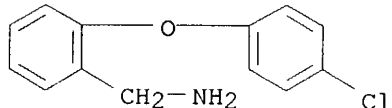


IT 146520-68-5P 146520-69-6P 146797-20-8P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
 (preparation of, as antidepressant)

RN 146520-68-5 HCAPLUS

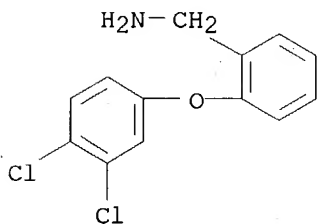
CN Benzenemethanamine, 2-(4-chlorophenoxy)-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

RN 146520-69-6 HCAPLUS

CN Benzenemethanamine, 2-(3,4-dichlorophenoxy)-, hydrochloride (9CI) (CA INDEX NAME)



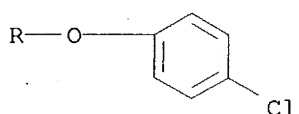
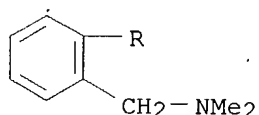
● HCl

RN 146797-20-8 HCAPLUS

CN Benzenemethanamine, 2-(4-chlorophenoxy)-N,N-dimethyl-, (2Z)-2-butenedioate

(1:1) (9CI) (CA INDEX NAME)

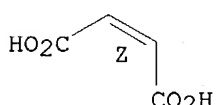
CM 1

CRN 146797-19-5
CMF C15 H16 Cl N O

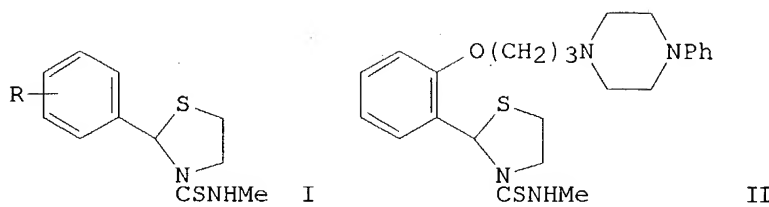
CM 2

CRN 110-16-7
CMF C4 H4 O4

Double bond geometry as shown.



L51 ANSWER 32 OF 44 HCPLUS COPYRIGHT 2004 ACS on STN
 ACCESSION NUMBER: 1988:75271 HCPLUS
 DOCUMENT NUMBER: 108:75271
 TITLE: Synthesis of 2-phenylthiazolidine derivatives as cardiotonic agents. I. 2-Phenylthiazolidine-3-thiocarboxamides
 AUTHOR(S): Nate, Hiroyuki; Sekine, Yasuo; Honma, Yasushi; Nakai, Hideo; Wada, Hiroshi; Takeda, Mikio; Yabana, Hideo; Nagao, Taku
 CORPORATE SOURCE: Org. Chem. Res. Lab., Tanabe Seiyaku Co., Ltd., Toda, 335, Japan
 SOURCE: Chemical & Pharmaceutical Bulletin (1987), 35(5), 1953-68
 DOCUMENT TYPE: Journal
 LANGUAGE: English
 OTHER SOURCE(S): CASREACT 108:75271
 GI



AB Novel 2-phenylthiazolidine-3-thiocarboxamides I ($R = 2$ -, 3-, 4-Cl, Me, OMe; 2-Me, 2-Et, 2-Pr, 2-Bu, 2-CH₂NMe₂, 2-NMe₂, 2-OH, 2-OCH₂Ph, 2-OCH₂CO₂H, 2-NHCOMe, 2-phenylpiperazinopropoxy etc.) was synthesized and tested for pos. inotropic activity in the isolated guinea pig heart and in anesthetized dogs. Reaction of RC₆H₄CHO with cysteamine followed by treatment with isothiocyanates readily gave I. Structure-activity relationships were investigated by varying the structural parameters. I ($R = 2$ -Me, 2-OMe) exhibited significant pos. inotropic action, which was not blocked by propranolol. α -Alkoxyphenyl derivative II exhibited more potent and longer-lasting activity than amrinone without any significant effect on heart rate or blood pressure.

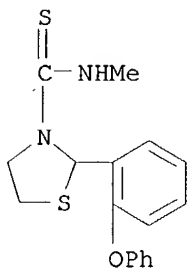
IT 112562-40-0P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation and inotropic activity of)

RN 112562-40-0 HCAPLUS

CN 3-Thiazolidinecarbothioamide, N-methyl-2-(2-phenoxyphenyl)- (9CI) (CA INDEX NAME)



L51 ANSWER 33 OF 44 HCAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: 1985:406319 HCAPLUS

DOCUMENT NUMBER: 103:6319

TITLE: A new method of synthesizing 8-10-membered heterocyclic systems condensed with two aromatic rings

Glinka, Ryszard; Piatowska, Elzbieta

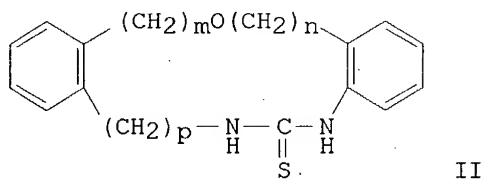
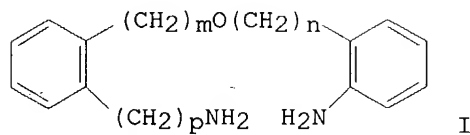
AUTHOR(S): Inst. Chem. Technol. Drugs, Sch. Med., Lodz, 90145, Pol.

CORPORATE SOURCE: Inst. Chem. Technol. Drugs, Sch. Med., Lodz, 90145, Pol.

SOURCE: Polish Journal of Chemistry (1984), 58(1-2-3), 259-62

CODEN: PJCHDQ; ISSN: 0137-5083

DOCUMENT TYPE: Journal
 LANGUAGE: English
 OTHER SOURCE(S): CASREACT 103:6319
 GI



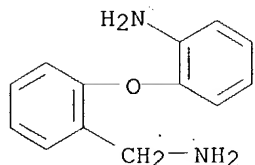
AB The diaminodiphenyl ether derivs I ($m = 0, 1, n = p = 0; m = n = 0, p = 1, m = n = 1, p = 0$) were treated with CS₂ in pyridine contg iodine to give the macrocyclic derivs II.

IT 30293-17-5

RL: RCT (Reactant); RACT (Reactant or reagent)
 (cyclization of, with carbon disulfide heterocyclic derivs. from)

RN 30293-17-5 HCPLUS

CN Benzenemethanamine, 2-(2-aminophenoxy)- (9CI) (CA INDEX NAME)



L51 ANSWER 34 OF 44 HCPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: 1984:191848 HCPLUS

DOCUMENT NUMBER: 100:191848

TITLE: Synthesis of N,N'-disubstituted derivatives of dibenzo[b,h]tetrahydro-1,4,6-oxadiazonine

AUTHOR(S): Glinka, Ryszard

CORPORATE SOURCE: Inst. Chem. Technol. Drugs, Sch. Med., Lodz, 90145, Pol.

SOURCE: Polish Journal of Chemistry (1983), Volume Date 1982, 56(7-8-9), 1139-44

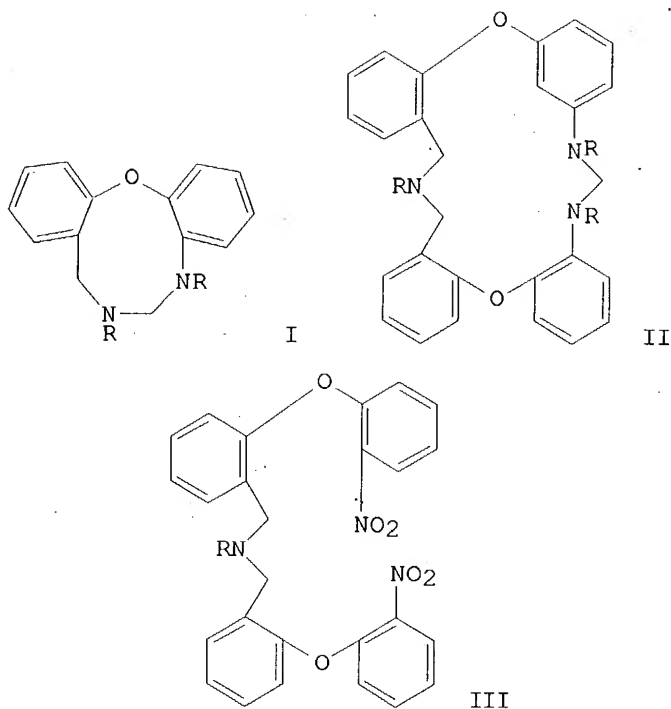
CODEN: PJCHDQ; ISSN: 0137-5083

DOCUMENT TYPE: Journal

LANGUAGE: English

OTHER SOURCE(S): CASREACT 100:191848

GI



AB The dibenzoxadiazonine I ($R = p\text{-MeC}_6\text{H}_4\text{SO}_2$ throughout) was prepared in 7 steps from $\text{o-MeC}_6\text{H}_4\text{OC}_6\text{H}_4\text{NO}_2\text{-o}$ via cyclization. of $\text{o-(RNNaCH}_2\text{)C}_6\text{H}_4\text{OC}_6\text{H}_4\text{(NNaR)-o}$ with BrCH_2Br . The tetrabenzotriazocyclohexadecane II was similarly prepared from the diether III in 3 steps.

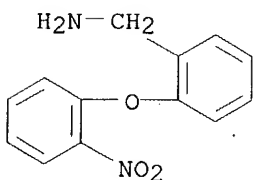
IT **89914-08-9P**

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and tosylation of)

RN 89914-08-9 HCPLUS

CN Benzenemethanamine, 2-(2-nitrophenoxy)-, monohydrochloride (9CI) (CA INDEX NAME)

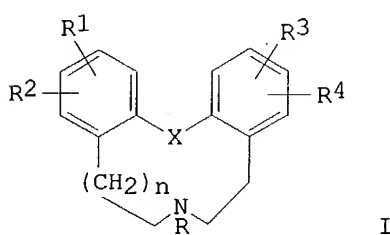


● HCl

L51 ANSWER 35 OF 44 HCAPLUS COPYRIGHT 2004 ACS on STN
 ACCESSION NUMBER: 1982:616253 HCAPLUS
 DOCUMENT NUMBER: 97:216253
 TITLE: Tricyclic compounds
 PATENT ASSIGNEE(S): AKZO N. V. , Neth.
 SOURCE: Neth. Appl., 16 pp.
 CODEN: NAXXAN
 DOCUMENT TYPE: Patent
 LANGUAGE: Dutch
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
NL 8006955	A	19820716	NL 1980-6955	19801222
WO 8202199	A1	19820708	WO 1981-EP195	19811212
W: AU, DK, FI, HU, JP				
AU 8279341	A1	19820720	AU 1982-79341	19811212
AU 547599	B2	19851024		
EP 57777	A1	19820818	EP 1981-201362	19811212
EP 57777	B1	19841010		
R: AT, BE, CH, DE, FR, GB, IT, LU, NL, SE				
JP 57502258	T2	19821223	JP 1982-500173	19811212
JP 02014350	B4	19900406		
HU 28460	O	19831228	HU 1982-110	19811212
HU 185195	B	19841228		
AT 9798	E	19841015	AT 1981-201362	19811212
US 4374133	A	19830215	US 1981-331303	19811216
ZA 8108782	A	19821124	ZA 1981-8782	19811218
ES 508218	A1	19830216	ES 1981-508218	19811221
CA 1174236	A1	19840911	CA 1981-392881	19811222
FI 74957	B	19871231	FI 1982-1944	19820601
FI 74957	C	19880411		
DK 8202724	A	19820708	DK 1982-2724	19820617
DK 151332	B	19871123		
DK 151332	C	19880606		
PRIORITY APPLN. INFO.:			NL 1980-6955	19801222
			EP 1981-201362	19811212
			WO 1981-EP195	19811212

GI



AB Dibenzoazacycloalkanes I ($X = O, S, NR_5$; $R = H, alkyl, aralkyl, hydroxyalkyl, acyloxyalkyl; R_1-R_4 = H, OH, halogen, cyano, alkyl, alkoxy, alkylthio, OCH_2O, CF_3, acyloxy; R_5 = H, alkyl; n = 0-2$) were prepared for use as tranquilizers (no data). Thus $O(C_6H_4CH_2CO_2H-2)_2$ was reduced to the diol, brominated, and treated with $PhCH_2NH_2$ to give I ($R = CH_2Ph, R_1-R_4 = H, X = O, n = 1$) which was debenzylated on Pd-C.

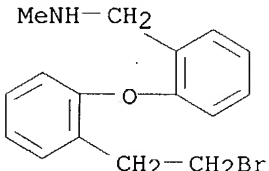
IT 83507-05-5P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and cyclization of)

RN 83507-05-5 HCAPLUS

CN Benzenemethanamine, 2-[2-(2-bromoethyl)phenoxy]-N-methyl- (9CI) (CA INDEX NAME)



L51 ANSWER 36 OF 44 HCAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: 1982:455432 HCAPLUS

DOCUMENT NUMBER: 97:55432

TITLE: Non-steroidal antiinflammatory agents. 7.

Methanesulfonanilides. II

AUTHOR(S): Schroeder, Eberhard; Lehmann, Manfred; Rufer, Clemens; Boettcher, Irmgard

CORPORATE SOURCE: Forschungslab., Schering A.-G., Berlin, 1000/65, Fed. Rep. Ger.

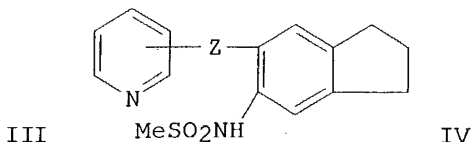
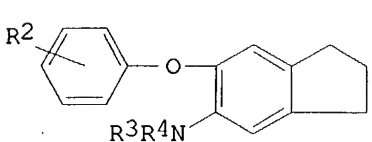
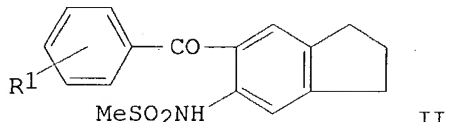
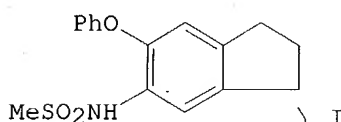
SOURCE: European Journal of Medicinal Chemistry (1982), 17(2), 165-72

DOCUMENT TYPE: CODEN: EJMCA5; ISSN: 0009-4374

LANGUAGE: Journal

OTHER SOURCE(S): German

GI OTHER SOURCE(S): CASREACT 97:55432



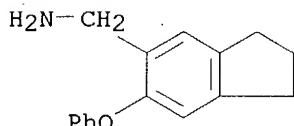
AB Analogs of the potent antiinflammatory sulfonanilide I were prepared and tested to determine structure-activity relationships. Of the compds. obtained by replacing the O atom of the PhO group by a direct bond, (substituted) methylene, or carbonyl group, only II (R1 = H, 2-Cl, 2-, 3-, 4-F) from the last procedure were active. Modifying the MeSO₂NH moiety caused loss of activity, but chloro- and fluoromethanesulfonates III (R2 = R4 = H, R3 = ClCH₂SO₂; R2 = H, 4-Cl, 4-F, R3 = F₃CSO₂, R4 = H) and Ac derivative III (R2 = H, R3 = MeSO₂, R4 = Ac) were active. The Ph ring was replaced with cyclohexyl, naphthyl, and heterocyclic groups and only pyridyl analogs IV (2-, 4-pyridyl, Z = S; 3-pyridyl, Z = O) showed antiinflammatory activity comparable to that of I.

IT 82472-01-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
(preparation and mesylation of)

RN 82472-01-3 HCPLUS

CN 1H-Indene-5-methanamine, 2,3-dihydro-6-phenoxy- (9CI) (CA INDEX NAME)



L51 ANSWER 37 OF 44 HCPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER:

1979:6244 HCPLUS

DOCUMENT NUMBER:

90:6244

TITLE:

Pyrrolidine derivatives

INVENTOR(S):

Beregi, Laszlo; Hugon, Pierre; Duhault, Jacques;
Boulanger, Michelle

PATENT ASSIGNEE(S):

Science Union et Cie., Societe Francaise de Recherche
Medicale, Fr.

SOURCE:

Ger. Offen., 21 pp.

CODEN: GWXXBX

DOCUMENT TYPE:

Patent

LANGUAGE:

German

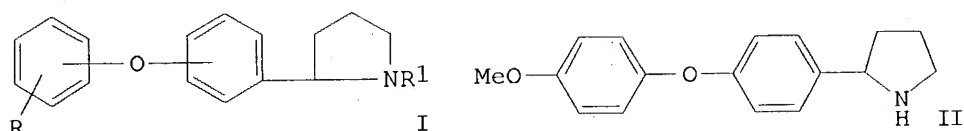
FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 2807623	A1	19780831	DE 1978-2807623	19780222
DE 2807623	C2	19820701		
GB 1561411	A	19800220	GB 1977-8096	19770225
US 4161529	A	19790717	US 1978-877601	19780214
CA 1101872	A1	19810526	CA 1978-297370	19780221
FR 2381753	A1	19780922	FR 1978-5005	19780222
FR 2381753	B1	19790713		
AT 7801289	A	19800715	AT 1978-1289	19780222
AU 7833552	A1	19790830	AU 1978-33552	19780223
BE 864312	A1	19780824	BE 1978-185474	19780224

NL 7802069	A	19780829	NL 1978-2069	19780224
JP 53105475	A2	19780913	JP 1978-20820	19780224
JP 55016577	B4	19800502		
ES 467286	A1	19781016	ES 1978-467286	19780224
DD 134088	C	19790207	DD 1978-203847	19780224
IN 147858	A	19800719	IN 1978-DE148	19780224
CH 631162	A	19820730	CH 1978-2058	19780224
ZA 7801100	A	19790131	ZA 1978-1100	19781218
PRIORITY APPLN. INFO.:			GB 1977-8096	19770225

GI



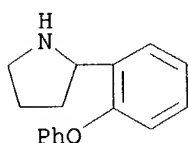
AB The phenoxyphenylpyrrolidines I [R = H, halogen, alkyl, alkoxy, CF₃; R₁ = H, C₁-4 aliphatic group, HO(CH₂)₂, HO(CH₂)₃, HO₂CCH₂] and their salts were prepared for use in regulation of lipid metabolism (no data). Thus, 4-MeOC₆H₄OC₆H₄COCl reacted with valine Me ester to give 4-(4-MeOC₆H₄O)C₆H₄CONHCH(CO₂Me)CHMe₂, which was saponified and treated successively with Ac₂O, CH₂:CHCN, and NaOH to give 4-(4-MeOC₆H₄O)C₆H₄COCH₂CH₂CN. Hydrogenative cyclization of this compound gave II.

IT 68548-77-6P 68548-78-7P

RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of)

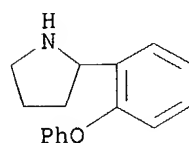
RN 68548-77-6 HCPLUS

CN Pyrrolidine, 2-(2-phenoxyphenyl)- (9CI) (CA INDEX NAME)



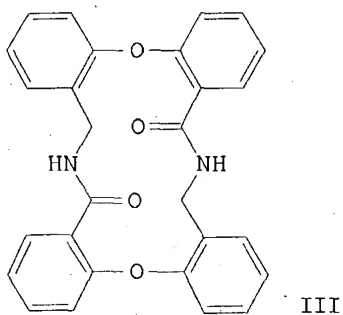
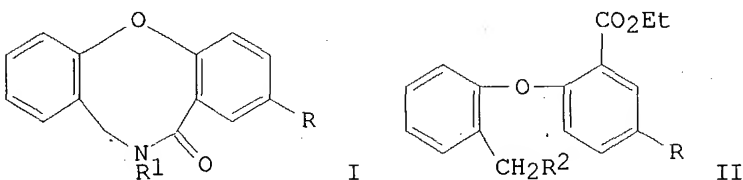
RN 68548-78-7 HCPLUS

CN Pyrrolidine, 2-(2-phenoxyphenyl)-, hydrochloride (9CI) (CA INDEX NAME)



●.HCl

L51 ANSWER 38 OF 44 HCPLUS COPYRIGHT 2004 ACS on STN
 ACCESSION NUMBER: 1976:164745 HCPLUS
 DOCUMENT NUMBER: 84:164745
 TITLE: Synthesis of 6,7-dihydrodibenz[b,g][1,5-oxazocin-5-ones
 ones
 AUTHOR(S): Lieb, Folker; Eiter, Karl
 CORPORATE SOURCE: Wiss. Hauptlab., Bayer A.-G., Leverkusen, Fed. Rep.
 Ger.
 SOURCE: Justus Liebigs Annalen der Chemie (1976), (2), 203-7
 DOCUMENT TYPE: CODEN: JLACBF; ISSN: 0075-4617
 LANGUAGE: Journal
 OTHER SOURCE(S): German
 CASREACT 84:164745
 GI



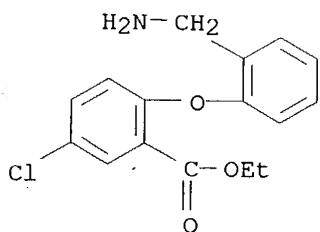
AB Dibenzoxazocinones I ($R = H$, $R1 = H$) were prepared by brominating II ($R2 = H$), treating II ($R2 = Br$) with K phthalimide, hydrazinolysis of II ($R2 =$ phthalimido), and cyclization of II ($R2 = NH_2$) with base. I ($R1 = H$) were aminoalkylated to give I ($R1 = CH_2CH_2NMe_2$). The dioxadiazacyclohexadecenedione III was prepared by treating 2-R₃OCH₂C₆H₄OCH₂C₆H₄CN-2 ($R3 = H$, Et) with H₂SO₄.

IT 59167-58-7P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
 (preparation and cyclization of)

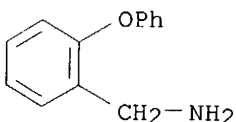
RN 59167-58-7 HCPLUS

CN Benzoic acid, 2-[2-(aminomethyl)phenoxy]-5-chloro-, ethyl ester,
 hydrochloride (9CI) (CA INDEX NAME)



● HCl

L51 ANSWER 39 OF 44 HCPLUS COPYRIGHT 2004 ACS on STN
 ACCESSION NUMBER: 1975:118784 HCPLUS
 DOCUMENT NUMBER: 82:118784
 TITLE: Antiarrhythmic agents. 2-, 3-, and 4-Substituted benzylamines
 AUTHOR(S): Remy, David C.; Van Saun, William A., Jr.; Engelhardt, Edward L.; Torchiana, Mary L.; Stone, Clement A.
 CORPORATE SOURCE: Merck Sharp and Dohme Res. Lab., Rahway, NJ, USA
 SOURCE: Journal of Medicinal Chemistry (1975), 18(2), 142-8
 CODEN: JMCMAR; ISSN: 0022-2623
 DOCUMENT TYPE: Journal
 LANGUAGE: English
 GI For diagram(s), see printed CA Issue.
 AB Of 29 title compds. prepared and tested for antiarrhythmic activity in dogs, 2-(*p*-methoxyphenylethylnyl)benzylamine-HCl (I-HCl) [54737-55-2], α,α -dimethyl-4-(phenylethylnyl)benzylamine-HCl (II-HCl) [38135-42-1], and α,α -dimethyl-4-phenethylbenzylamine-HCl (III-HCl) [38135-43-2] showed good activity. Many of the compds. are more active antiarrhythmic agents than quinidine [56-54-2] tested under similar conditions. Structure-activity relations are discussed.
 IT 31963-35-6P
 RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
 (preparation and antiarrhythmic activity of)
 RN 31963-35-6 HCPLUS
 CN Benzenemethanamine, 2-phenoxy-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

L51 ANSWER 40 OF 44 HCAPLUS COPYRIGHT 2004 ACS on STN
 ACCESSION NUMBER: 1971:141251 HCAPLUS
 DOCUMENT NUMBER: 74:141251
 TITLE: Antiarrhythmic o-phenoxybenzylamine
 INVENTOR(S): Remy, David C.
 PATENT ASSIGNEE(S): Merck and Co., Inc.
 SOURCE: Ger. Offen., 26 pp.
 CODEN: GWXXBX
 DOCUMENT TYPE: Patent
 LANGUAGE: German
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 2047677	A	19710408	DE 1970-2047677	19700928
NL 7013566	A	19710331	NL 1970-13566	19700914
GB 1287497	A	19720831	GB 1970-1287497	19700928
FR 2070095	A5	19710910	FR 1970-35195	19700929
FR 2070095	B1	19750418		

PRIORITY APPLN. INFO.: US 1969-862076 19690929

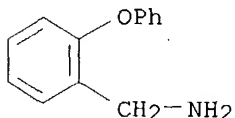
AB The title compound was prepared by reaction of o-BrC₆H₄CN with PhONa in MeOH in the presence of Cu to give o-NCC₆H₄OPh which was reduced with LiAlH₄ in Et₂O.

IT 31963-35-6P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of)

RN 31963-35-6 HCAPLUS

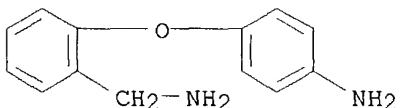
CN Benzenemethanamine, 2-phenoxy-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

L51 ANSWER 41 OF 44 HCAPLUS COPYRIGHT 2004 ACS on STN
 ACCESSION NUMBER: 1971:3381 HCAPLUS
 DOCUMENT NUMBER: 74:3381
 TITLE: Nucleophilic aromatic substitution by aminophenoxy ions
 AUTHOR(S): Schramm, Juergen; Radlmann, Eduard; Lohwasser, Hermann; Nischk, Guenther
 CORPORATE SOURCE: Org.-Wiss. Lab., Farbenfabriken Bayer A.-G., Dormagen/Rhein, Fed. Rep. Ger.
 SOURCE: Justus Liebigs Annalen der Chemie (1970), 740, 169-79
 DOCUMENT TYPE: Journal
 CODEN: JLACBF; ISSN: 0075-4617

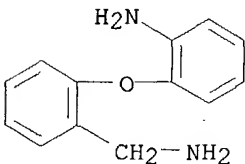
LANGUAGE: German
 OTHER SOURCE(S): CASREACT 74:3381
 AB Aromatic RC1 activated by o-or (and) p-NO₂ reacted with x-H₂NC₆H₄OK (I) in Me₂SO at room temperature to give x-H₂NC₆H₄OR (II) (x = o-, m-, or p-position,
 R = C₆H₄NO₂-p or o-, C₆H₃(NO₂)Cl-4,2, C₆H₃(NO₂)CF₃-2,4, C₆H₃-(NO₂)₂-2,4], in some cases addnl. in minor amts. 2,4-R₁(O₂N)-C₆H₃OC₆H₄[NHC₆H₃(NO₂)R₁-4,2]-4 (R₁ = H, Cl, NO₂), and 4-HOC₆H₄NHC₆H₃(NO₂)R₁-4,2, if R₁ = NO₂. 6,3- and 3,4-Cl-(H₂N)C₆H₃OK reacted similarly to give the II-analogs. II were hydrogenated over Raney Ni to give the corresponding diamines. Similarly, o- and p-ClC₆H₄CN and I gave the corresponding H₂NC₆H₄OC₆H₄CN, which were hydrogenated as above. Dichloroaryl and -heteroaryl compds. and I gave the corresponding H₂NC₆H₄OQOC₆H₄NH₂ (Q = 3,4-pyridazinylene, 2,4-pyrimidinylene, 1,4-phthalazinylene, 1,3-C₆H₃NO₂-6, 1,3-C₆H₂(NO₂)Cl-6,4, or 4-C₆H₄SO₂C₆H₄-4).
 IT 30202-94-9P 30203-00-0P 30203-01-1P
 30293-17-5P
 RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of)
 RN 30202-94-9 HCPLUS
 CN Benzylamine, o-(p-aminophenoxy)- (8CI) (CA INDEX NAME)



RN 30203-00-0 HCPLUS
 CN Benzylamine, o-(o-aminophenoxy)-, monopicrate (8CI) (CA INDEX NAME)

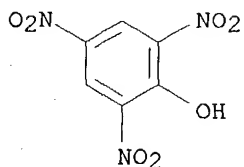
CM 1

CRN 30293-17-5
CMF C₁₃ H₁₄ N₂ O

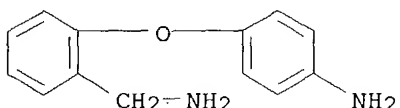


CM 2

CRN 88-89-1
CMF C₆ H₃ N₃ O₇

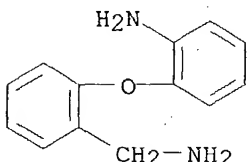


RN 30203-01-1 HCPLUS
 CN Benzylamine, o-(p-aminophenoxy)-, dihydrochloride (8CI) (CA INDEX NAME)



●2 HCl

RN 30293-17-5 HCPLUS
 CN Benzenemethanamine, 2-(2-aminophenoxy)- (9CI) (CA INDEX NAME)



L51 ANSWER 42 OF 44 HCPLUS COPYRIGHT 2004 ACS on STN
 ACCESSION NUMBER: 1970:466188 HCPLUS
 DOCUMENT NUMBER: 73:66188
 TITLE: Macro chelate rings. III. Syntheses and configurations of complexes of new ligands, 4,4'-dimethyl-2,2'-bis(salicylideneaminomethyl)diphenyl ether and 4,4'-dimethyl-2,2'-bis(salicylideneaminomethyl)diphenylamine
 AUTHOR(S): Okawa, Hisashi; Koyama, Hiroyuki; Inazu, Takahiko; Yoshino, Tamotsu
 CORPORATE SOURCE: Fac. Sci., Kyushu Univ., Fukuoka, Japan
 SOURCE: Bulletin of the Chemical Society of Japan (1970), 43(6), 1729-33
 CODEN: BCSJA8; ISSN: 0009-2673
 DOCUMENT TYPE: Journal
 LANGUAGE: English
 GI For diagram(s), see printed CA Issue.
 AB Two quadridentate chelating agents, 4,4'-dimethyl-2,2'-bis(salicylideneaminomethyl)diphenyl ether (I) and 4,4'-dimethyl-2,2'-

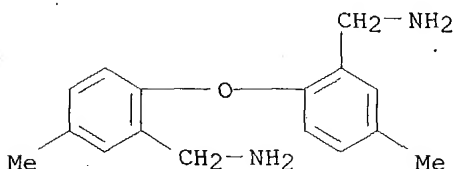
bis(salicylideneaminomethyl)diphenylamine (II), which may form metal complexes with a ten-membered chelate ring, were synthesized and their Co(II), Ni(II), Cu(II), and Zn(II) complexes prepared. The configurations around the central divalent metal ion were explored on the basis of the absorption of the ligand field in a solution. A tetrahedral configuration for Co(II) and Zn(II) complexes, and a pseudotetrahedral configuration for the Ni(II) and Cu complexes, were concluded.

IT 27996-12-9P

RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of)

RN 27996-12-9 HCPLUS

CN Benzylamine, 2,2'-oxybis[5-methyl-, dihydrochloride (8CI) (CA INDEX NAME)



●2 HCl

L51 ANSWER 43 OF 44 HCPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: 1961:118547 HCPLUS

DOCUMENT NUMBER: 55:118547

ORIGINAL REFERENCE NO.: 55:22321f-i,22322a-b

TITLE: The synthesis of esters of some amino acids having pharmacological importance. I. The synthesis of esters of piperidino carboxylic acids

AUTHOR(S): Matkovics, Bela; Foldeak, Sandor; Porszasz, Janos; Sipos, Gyorgy

CORPORATE SOURCE: Tudomanyegyetem, Szeged, Hung.

SOURCE: Acta Pharmaceutica Hungarica (1961), 31, 113-21

CODEN: APHGAO; ISSN: 0001-6659

DOCUMENT TYPE: Journal

LANGUAGE: Hungarian

AB RCH₂CO₂R' (I), RCH₂CH₂CO₂R' (II), BzOCH₂CH₂R (III), and AcOCHMeCH₂R (IV) were prepared. I were prepared by condensing ClCH₂CO₂R' with a secondary amine, II by boiling ClCH₂CH₂CO₂R' with the amine, and III by the reaction of an amino alc. with BzCl. The following I were obtained (R, R', b.p.°/mm., m.p. of picrate, m.p. of HCl salt, and m.p. of methiodide are given): piperidino, Me, 69°/5, 115°, 214°, 163-4°; piperidino, Et, 68°/1, 122°, 117-17.5°, 160-60.3°; piperidino, Bu, 100-1°/4, 85°, -, 178°; piperidino, PhCH₂, 134-5°/1, 137°, 133°, 91-6°; morpholino, Me, 77°/2, 143°, 150.5°, 147.5°; morpholino, Et, 86-7°/4, 163°, 181°, 132-3°; morpholino, Bu, 105.5-106°/3, -, 127-9°, 95-6°; morpholino, PhCH₂, 164-5°/5, 143°, 149°, -; pyrrolidino, Me, 72-3°/8, 104°, -, 153°; pyrrolidino, Et,

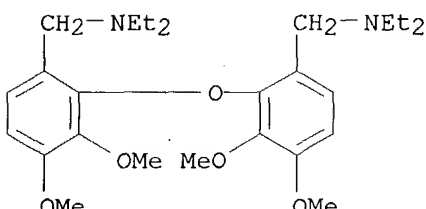
59-60°/2, 119.5°, 133-3.5°, -; pyrrolidino, Bu,
 81-2°/3, 109.5°, -, -; pyrrolidino, PhCH₂, 134-5°/1,
 159-60°, 139-40°, 156°. The following II were prepared
 (data as above): piperidino, Me, 72°/2, 164°, 189°,
 147-8°; piperidino, Et, 102-3°/5, 131.5°,
 169°, -; piperidino, Bu, 124-5°/6, 108-9°,
 164.7°, -; piperidino, PhCH₂, 149-50°/1, 113°,
 193.5°, -; piperidino, Ph, 114-20°/3, -, 192-5°, -;
 piperidino, CPh₃, 171°/1, -, 214°, -; morpholino, Me,
 82°/2, 129°, 203°, 151°; morpholino, Et,
 108°/6, 108°, 188-9°, -; morpholino, Bu,
 131-2°/6, 150°, 173°, 115°; morpholino, PhCH₂,
 154°/1, 125°, 189-90°, -; pyrrolidino, Me,
 76°/5, 147°, 128°, 166°; pyrrolidino, Et,
 85°/6, 114°, 146°, -; pyrrolidino, Bu,
 106-8°/5, 97°, 74-5°, 115°; pyrrolidino,
 PhCH₂, 145-6°/3, 102°, 152°, 154°. IV (R
 =pyrrolidino) (V), b3 75°, picrate m. 111-12°, gave a
 hygroscopic HCl salt. III (R = piperidino) b2 141°; HCl salt m.
 184°; methiodide m. 141.5°. The action of the compds. on
 blood pressure and on respiration was given. II (R = N-piperidino, R' =
 CPh₃) and V had strong antinicotinic action. The effect of the piperidino
 and pyrrolidino propionates was increased by quaternization.

IT 103044-26-4, Veratrylamine, 2,2'-oxybis[N,N-diethyl-

(preparation of)

RN 103044-26-4 HCPLUS

CN Veratrylamine, 2,2'-oxybis[N,N-diethyl- (6CI) (CA INDEX NAME)



L51 ANSWER 44 OF 44 HCPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: 1957:71530 HCPLUS

DOCUMENT NUMBER: 51:71530

ORIGINAL REFERENCE NO.: 51:12942c-i,12943a-i,12944a-c

TITLE: Alkaloid studies. XVII. The structure of the cactus
 alkaloid pilocereine

AUTHOR(S): Djerassi, Carl; Figgord, S. K.; Bobbitt, J. M.;
 Markley, F. X.

CORPORATE SOURCE: Wayne State Univ., Detroit, MI

SOURCE: Journal of the American Chemical Society (1957), 79,
 2203-10

CODEN: JACSAT; ISSN: 0002-7863

DOCUMENT TYPE: Journal

LANGUAGE: Unavailable

GI For diagram(s), see printed CA Issue.

AB cf. C.A. 51, 8118d. Structure I (R = CH₂CHMe₂) was elucidated for the

cactus alkaloid pilocereine. I (8.5 g.) in 200 cc. MeOH-280 cc. Et₂O treated 6 days at 0° with 2.2 g. distilled CH₂N₂, the mixture treated with an addnl. 2.2 g. CH₂N₂, kept 3 days at 0°, and evaporated and the residue recrystd. from hexane yielded 6.5 g. Me ether (II) of I, m. 92-105°, resolidified and m. 153-5° (all m.ps. were determined on a Kofler block). II, m. 153-5° (from EtOAc), was transformed to a 2nd crystalline form, m. 133-5°; the transformation was reversed by recrystn. from hexane. I (3.0 g.) in 100 cc. absolute EtOH treated with 3.6 g. MeCHN₂ in 150 cc. Et₂O, kept 24 hrs. at room temperature, treated with an addnl. 3.6 g. MeCHN₂, refrigerated 6 days, and evaporated yielded 2.07 g. Et ether (III) of I, m. 90-5° and 152-3° (from hexane); 2nd crop, 0.32 g. Amberlite IRA-400 (HCl) (200 g.) treated with 500 cc. 50% aqueous NaOH, 2 l. H₂O, and finally 250 g. NaHCO₃ in saturated aqueous solution and

washed with 12-16 l. H₂O gave the bicarbonate salt IRA-400-HCO₃ which was stored under distilled H₂O. Styphnates and picrates in EtOH or Me₂CO containing

about 5% H₂O passed dropwise over a column of IRA-400-HCO₃, the column washed with 2 vols. 10% aqueous Me₂CO, the Me₂CO removed in vacuo, acid added, the aqueous solution washed with Et₂O and basified with NH₄OH, and the base isolated with Et₂O gave the corresponding free amines. II (2.5 g.) in 100 cc. 10% H₂SO₄ made just alkaline with 2N NaOH, treated dropwise at room temperature

with 250 cc. 2% aqueous KMnO₄, allowed to stand overnight, acidified with H₂SO₄, and extracted continuously with Et₂O, the residue from the extract treated

with SOC₁₂ and then PhNH₂, and the product chromatographed yielded 35 mg. iso-PrCONHPh and 10 mg. iso-BuCONHPh. I (5.0 g.) in 200 cc. dry Et₂O added slowly with stirring to 1.5 l. liquid NH₃ at -60° during 5 hrs., the mixture warmed during 3 hrs. to -30°, treated cautiously with NH₄Cl and evaporated overnight, the residue partitioned between Et₂O and 3% aqueous NaOH, the alkaline layer acidified with 40% H₂SO₄, washed with Et₂O, basified with concentrated NH₄OH, and extracted with Et₂O, and the extract evaporated gave

2.46 g. phenolic basic oil (IV); the original Et₂O layer extracted with 10% HCl, dried, and evaporated left only a small amount of nonphenolic, nonbasic oil

which was discarded; the acid extract basified with NH₄OH and extracted with Et₂O

gave 2.40 g. nonphenolic, basic, glassy material (V). V consisted mainly of isopilocereine (VI); dipicrate, m. 235-7° (from Me₂CO). VI dipicrate (3.5 g.) treated with LiOH and the resulting free base treated with CH₂N₂ in Et₂O-MeOH yielded 55% Me ether (VII) of VI, b0.005 180-90° (evaporatively distilled). In 1 run, a 75-mg. aliquot of V treated with 40 mg. picric acid yielded 70 mg. 1-isobutyl-2-methyl-6-methoxy-1,2,3,4-tetrahydroisoquinoline (VIII) picrate, m. 150-1° (from MeOH). IV (0.26 g.) treated 6 days at 0° with CH₂N₂ in Et₂O containing a small amount of MeOH and evaporated, the residue extracted with Et₂O and

washed with 3% aqueous NaOH, and the resulting oil (0.2 g.) chromatographed on 9 g. Al₂O₃ gave 0.155 g. 7-MeO derivative (IX) of VIII, nD₂₅ 1.5284; styphnate, m. 212-13°; picrate, m. 184-5°. I (5 g.) in 1.5 l. dry NH₃ treated at -30° with 6 g. K, and the mixture worked up in the usual manner gave 1.79 g. V and 2.68 g. IV; the IV dissolved in Et₂O, dried, and concentrated yielded 1.45 g. demethylisopilocereine (X), m. 177.5-78°. X (100 mg.) treated 2 days at 0° with excess CH₂N₂ in Et₂O and evaporatively distilled yielded 81 mg. glass, the infrared

spectrum of which closely resembled that of VI; treatment with picric acid gave a small amount of VI picrate. X (210 mg.) in Et₂O-MeOH treated 7 days with CH₂N₂ yielded 120 mg. VII. IV (300 mg.) treated 7 days at room temperature

with 0.84 g. MeCHN₂ in Et₂O, washed with alkali, and treated with picric acid gave the picrate of the 7-EtO derivative of VIII, m. 151.5-2.5°. Natural IX (2.2 g.) oxidized with KMnO₄ yielded 310 mg. m-hemipinic acid, characterized as the di-Me ester, m. 89.5-90°; iso-PrCO₂H and iso-BuCO₂H were identified as their anilides. IX (2.47 g.) and 10 cc. MeI kept overnight at room temperature, the resulting methiodide (5.17 g.) dissolved

in a small amount of H₂O, added to 120 cc. 50% aqueous KOH, and refluxed 2 hrs.,

and the product isolated in the usual manner yielded 2.05 g. 2,4,5-[iso-Bu(Me₂N)CH](MeO)C₆H₂CH:CH₂ (XI), oil. XI (185 mg.) in glacial AcOH ozonized 0.5 hr. at 15° and steam distilled into dimedon in MeOH, and the mixture kept 24 hrs. at 0° gave 39 mg. CH₂O derivative, m. 193-5°. XI (1.87 g.) in MeOH hydrogenated 1 hr. over 5% Pd-C yielded the 1-Et analog (XII) of XI. XII converted to the methiodide (3.94 g.) and boiled with 50% aqueous KOH yielded 1.06 g. neutral N-free oil, apparently 3,4,5-Et(MeO)C₆H₂CH:CHCHMe₂; a 90-mg. portion ozonized and steam distilled into acidified aqueous 2,4-(O₂N)C₆H₃NHNH₂, extracted with C₆H₆, and

chromatographed on Al₂O₃ yielded 20 mg. iso-PrCHO derivative, m. 181-2°. IX oxidized with KMnO₄ in the same manner as I gave iso-PrCO₂H and iso-BuCO₂H. VII (104 mg.) in C₆H₆ treated 4.5 hrs. with 1 cc. MeI gave 153 mg. VII.2MeI, m. 191-4° (from hexane-Me₂CO). VII.2MeI (150 mg.) in 5 cc. MeOH and 20 cc. H₂O passed 4 times over IRA-400-OH resin, the column washed with 20 cc. 50% aqueous MeOH, and the residue from the eluates distilled yielded 89 mg. gummy methine, C₃₃H₅₀N₂O₄, b0.05, 170-5°; a 100-mg. sample ozonized in CHCl₃ at -60° gave 55 mg. CH₂O-dimedon derivative; a 500-mg. sample in EtOH hydrogenated 10 min. over Pd-C yielded 450 mg. reduced methine (XIII), b0.005 160° (bath temperature). XIII (130 mg.) in Et₂O treated with MeI, the dimethiodide (180 mg.) decomposed by the ion exchange resin method, the resulting neutral olefin (76 mg.), b0.005 160-80° ozonized in CHCl₃, at -60°, and the distillate passed into 2,4-(O₂N)C₆H₃NHNH₂ solution yielded 44% 2,4-(O₂N)₂C₆H₃NHN:CHCHMe₂ (XIIIa). II (2.56 g.) treated with MeI, the II.MeI (3.9 g.), m. 137-50° (decomposition), powdered, added to 100 cc. refluxing 40% aqueous NaOH, and refluxed 2.5 hrs., a 160-mg. portion of the resulting methine 4,2,-5-R(MeO)[CH(NMe₂)(CH₂CHMe₂)C₆H₂OCH(OME)₂[CH(NMe₂)(CH₂CHMe₂)]R-2,3,6,5 (XIV) (R = CH:CH₂) (2.0 g.) ozonized in AcOH, and the mixture steam distilled into 2,4-(O₂N)C₆H₃NHNH₂ gave only 47 mg. CH₂O derivative

XIV (R = CH:CH₂) (1.9 g.) in 50 cc. 95% EtOH hydrogenated over 300 mg. 10% Pd-C, and the crude product (1.85 g.) recrystd. from MeCN gave 0.92 g. XIV (R = Et), m. 101.5-3.5°. XIV (R = Et) (1.21 g.) subjected to a 2nd stage Hofmann degradation gave 0.45 g. Me₃N picrate, m. 206-10°, and 0.84 g. N-free degradation product which ozonized in EtOAc at -60° and worked up in the usual manner yielded only 3% XIIIa. XIV (R = Et) converted to the dimethiodide (1.72 g.) and subjected to a Hofmann degradation in the usual manner except that the compound was first dissolved in EtOH gave a substance, b0.005 155-70°, which appeared to be the di-CH(OEt)CH₂CHMe₂ analog (XV) of XIV (R = Et). II (1.98 g.) cleaved in the usual manner with 90 cc. Et₂O, 600 cc. liquid NH₃, and 2.5 g. K at -60° during 7 hrs. gave 1.30 g. nonphenolic basic and 0.67 g. phenolic basic fractions. The nonphenolic fractions dissolved in 20

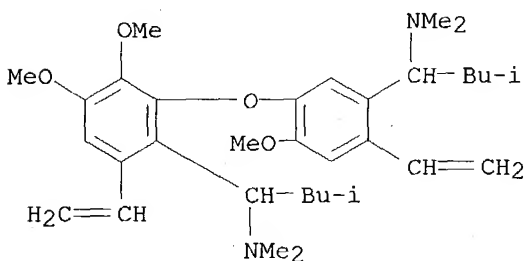
cc. hexane and chromatographed on 80 g. Al2O3 (deactivated with 2.4 cc. 10% AcOH), giving 114 fractions, and fractions 20-46 (hexane up to 1:1 hexane-C6H6) treated with alc. picric acid gave 0.53 g. picrate of VIII, m. 152-3°; fractions 47-83 (1:1 hexane-C6H6 to 99:1 C6H6-Et2O) treated with alc. picric acid gave 0.196 g. IX picrate, m. 183-5°. Fractions 100-12 (9:1 C6H6-Et2O) gave similarly 10% picrate of the 8-OH derivative (XVI) of IX, m. 150-5°. XVI (73 mg.) (from the picrate) treated 10 days at 0° with CH2N2 in Et2O-MeOH and the product treated with alc. picric acid yielded the picrate of the 8-MeO analog (XVII) of XVI, m. 132-4°. Fractions 112-14 (Et2O and 9:1 Et2O-MeOH) gave a picrate, m. unsharply above 210°, which may represent dimeric material. The phenolic cleavage product (0.67 g.) and CH2N2 in MeOH-Et2O refrigerated 8 days yielded 0.43 g. picrate of IX, m. 181-4°; the mother liquors transformed to the free amine by the ion exchange method and chromatographed on deactivated Al2O3 gave 0.164 g. oil which treated with picric acid yielded 0.175 g. picrate of XVII. III (2.04 g.) in 80 cc. Et2O and 600 cc. liquid NH3 treated at -60° with 3.3 g. K and the mixture worked up after 24 hrs. gave 1.30 g. nonphenolic basic and 0.51 g. phenolic basic fractions. The nonphenolic portion chromatographed in the usual manner gave 0.576 g. VIII picrate, m. 151-3°, 0.227 g. picrate of the 7-EtO analog (XVIII) of IX, m. 152-3°, and 0.244 g. picrate of the 8-OH derivative of XVIII, m. 153-4°. The phenolic portion (0.51 g.) methylated in the usual manner and treated with picric acid gave 0.356 g. picrate of IX, m. 183-5°.

IT 115606-43-4, Veratrylamine, 2-[5-(1-dimethylamino-3-methylbutyl)-2-methoxy-4-vinylphenoxy]- α -isobutyl-N,N-dimethyl-6-vinyl-

117272-08-9, Veratrylamine, 2-[5-(1-dimethylamino-3-methylbutyl)-4-ethyl-2-methoxyphenoxy]-6-ethyl- α -isobutyl-N,N-dimethyl- (preparation of)

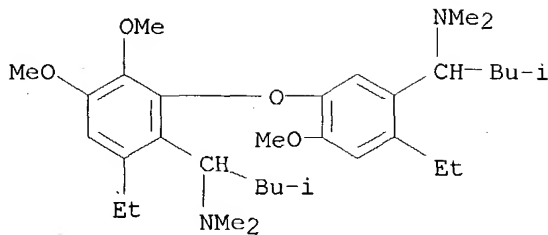
RN 115606-43-4 HCPLUS

CN Veratrylamine, 2-[5-(1-dimethylamino-3-methylbutyl)-2-methoxy-4-vinylphenoxy]- α -isobutyl-N,N-dimethyl-6-vinyl- (6CI) (CA INDEX NAME)



RN 117272-08-9 HCPLUS

CN Veratrylamine, 2-[5-(1-dimethylamino-3-methylbutyl)-4-ethyl-2-methoxyphenoxy]-6-ethyl- α -isobutyl-N,N-dimethyl- (6CI) (CA INDEX NAME)



=> b caold

FILE 'CAOLD' ENTERED AT 11:29:44 ON 11 FEB 2004
USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.
PLEASE SEE "HELP USAGETERMS" FOR DETAILS.
COPYRIGHT (C) 2004 AMERICAN CHEMICAL SOCIETY (ACS)

FILE COVERS 1907-1966

FILE LAST UPDATED: 01 May 1997 (19970501/UP)

This file contains CAS Registry Numbers for easy and accurate substance identification. Title keywords, authors, patent assignees, and patent information, e.g., patent numbers, are now searchable from 1907-1966. TIFF images of CA abstracts printed between 1907-1966 are available in the PAGE display formats.

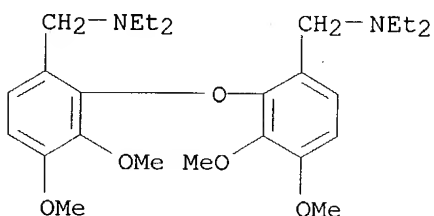
This file supports REGISTRY for direct browsing and searching of all substance data from the REGISTRY file. Enter HELP FIRST for more information.

=> d que 154 nos

L6	STR
L7	STR
L8	STR
L9 (6520) SEA FILE=REGISTRY SSS FUL L6
L10 (747) SEA FILE=REGISTRY SUB=L9 SSS FUL (L6 AND (L7 OR L8))
L11	STR
L12	STR
L13	STR
L14	STR
L15	544 SEA FILE=REGISTRY SUB=L10 SSS FUL (L6 AND L14 NOT (L11 OR L12 OR L13))
L17	STR
L20	STR
L46	STR
L47	STR
L48	STR
L50	273 SEA FILE=REGISTRY SUB=L15 SSS FUL (L17 AND (L20 OR L46 OR L47 OR L48))
L54	3 SEA FILE=CAOLD ABB=ON PLU=ON L50

=> d iall hitstr 1-3

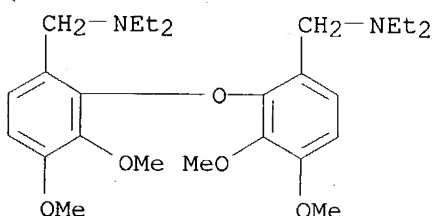
L54 ANSWER 1 OF 3 CAOLD COPYRIGHT 2004 ACS on STN
 ACCESSION NUMBER: CA55:22321e CAOLD
 TITLE: synthesis and halomethylation of bis(3,4-dimethoxyphenyl) ether-reaction of halomethyl derivs. with secondary amines and pyridines
 AUTHOR NAME: Matarasso-Tchiroukhine, Elisabeth
 INDEX TERM: 786-19-6 101432-12-6 101432-17-1 101744-16-5 102015-72-5
 102015-78-1 **103044-26-4** 108480-92-8 108480-93-9
116378-59-7 116572-03-3 116572-04-4 117042-78-1
 117042-83-8
 IT **103044-26-4** **116378-59-7**
 RN 103044-26-4 CAOLD
 CN Veratrylamine, 2,2'-oxybis[N,N-diethyl- (6CI) (CA INDEX NAME)



RN 116378-59-7 CAOLD
 CN Veratrylamine, 2,2'-oxybis[N,N-diethyl-, dipicrate (6CI) (CA INDEX NAME)

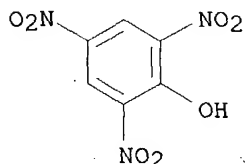
CM 1

CRN 103044-26-4
 CMF C26 H40 N2 O5



CM 2

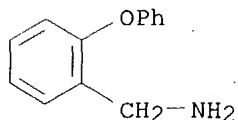
CRN 88-89-1
 CMF C6 H3 N3 O7



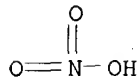
L54 ANSWER 2 OF 3 CAOLD COPYRIGHT 2004 ACS on STN
 ACCESSION NUMBER: CA52:9005i CAOLD
 TITLE: synthesis of guanidine compds. of diphenyl ether - (I), (II)
 synthesis of diphenyl ether aldehyde by the Sommelet
 reaction and expts. with methylguanidine derivs. (1)
 AUTHOR NAME: Ito, Genzo
 INDEX TERM: 67-36-7 500-78-7 3396-01-8 19434-34-5 22479-78-3
 36881-42-2 62248-88-8 76838-41-0 76839-21-9 78725-47-0
 82657-72-5 91955-44-1 92028-82-5 100394-74-9 100714-40-7
 100724-23-0 100956-00-1 100970-01-2 101091-62-7 101097-62-5
 101112-34-9 101868-83-1 102078-83-1 102309-41-1 105901-52-8
 105901-53-9 106272-17-7 107558-81-6 107622-81-1
107624-15-7 107771-82-4 107774-30-1 107776-32-9
 107921-04-0 107922-64-5 108125-78-6 108752-39-2 108758-94-7
 108800-18-6 108874-34-6 108923-35-9 108955-35-7 108955-88-0
 109017-27-8 109102-87-6 109125-85-1 109125-91-9 109125-92-0
 109556-73-2 109592-53-2 109592-68-9 110533-52-3 110974-03-3
 114281-51-5 119077-06-4 119438-84-5 123777-32-2 124111-89-3
 125644-23-7 132624-85-2

IT **107624-15-7**
 RN 107624-15-7 CAOLD
 CN Benzylamine, o-phenoxy-, nitrate (6CI) (CA INDEX NAME)

CM 1

CRN 107624-14-6
CMF C13 H13 N O

CM 2

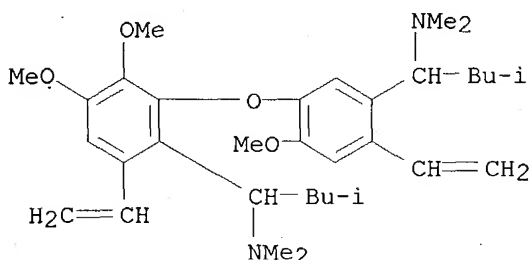
CRN 7697-37-2
CMF H N O3

L54 ANSWER 3 OF 3 CAOLD COPYRIGHT 2004 ACS on STN
 ACCESSION NUMBER: CA51:12942c CAOLD
 TITLE: alkaloid studies - (XVII) structure of the cactus alkaloid pilocereine
 AUTHOR NAME: Djerassi, Carl; Figidor, S. K.; Bobbitt, J. M.; Markley, F. X.
 INDEX TERM: 479-49-2 3300-36-5 3423-32-3 25181-15-1 91809-56-2
 93146-43-1 94326-14-4 95282-34-1 96506-44-4 101745-56-6
 101745-83-9 102032-23-5 102589-65-1 103404-48-4 105820-94-8
 111979-41-0 112045-14-4 115604-83-6 **115606-43-4**
 116571-90-5 **117272-08-9** 121233-16-7 124142-92-3
 124515-45-3

IT **115606-43-4 117272-08-9**

RN 115606-43-4 CAOLD

CN Veratrylamine, 2-[5-(1-dimethylamino-3-methylbutyl)-2-methoxy-4-vinylphenoxy]- α -isobutyl-N,N-dimethyl- (6CI) (CA INDEX NAME)



RN 117272-08-9 CAOLD

CN Veratrylamine, 2-[5-(1-dimethylamino-3-methylbutyl)-4-ethyl-2-methoxyphenoxy]-6-ethyl- α -isobutyl-N,N-dimethyl- (6CI) (CA INDEX NAME)

